Office européen des brevets



EP 0 371 998 B2

(12)

NEW EUROPEAN PATENT SPECIFICATION

- (45) Date of publication and mention of the opposition decision: 03.11.1999 Bulletin 1999/44
- (45) Mention of the grant of the patent: 02.03.1994 Bulletin 1994/09
- (21) Application number; 88907510.7
- (22) Date of filing: 25.07.1988

UTILISATION

- (51) Int Ci.6: C12N 15/13, C12N 15/70, C12N 15/85, C12N 1/21, C12N 5/10
- (86) International application number: PCT/US88/02514
- (87) International publication number, WO 89/00999 (09.02.1989 Gazette 1989/04)
- (54) MODULAR ASSEMBLY OF ANTIBODY GENES, ANTIBODIES PREPARED THEREBY AND USE MODULÄRER ZUSAMMENBAU VON ANTIKÖRPERGENEN, DADURCH HERGESTELLTE ANTIKÖRPER UND DEREN ANWENDUNG ENSEMBLE MODULAIRE DE GENES D'ANTICORPS, ANTICORPS AINSI PREPARES ET
- (84) Designated Contracting States: AT BE CHIDE FRIGBIT LI LU NL SE
- (30) Priority: 24.07.1987 US 77528
- (43) Date of publication of application: 13.06.1990 Bulletin 1990/24
- (60) Divisional application: 93100041.8 / 0 550 400
- (73) Proprietor. XOMA Corporation Berkeley California 94710-2737 (US)
- (72) Inventors:
 - ROBINSON, Randy, R. Los Angeles, CA 90045 (US)
 - . LIU. Aivin. Y. Oceanside, CA 92056 (US)
 - . HORWITZ, Arnold, H.
 - Los Angeles, CA 90045 (US)
 - · WALL, Randolph
 - Sherman Oaks, CA 91423 (US) BETTER, Marc
 - Los Angeles, CA 90046 (US)
- (74) Representative: Ritter, Stephen David et al

Mathys & Squire 100 Grays Inn Road London WC1X 8AL (GB)

(56) References cited: EP-A- 0 125 023 EP-A- 0 173 494 EP-A- 0 234 592

EP-A1- 0 120 694 EP-A1- 0 126 338 WO-A-86/01533 GB-A- 2 137 631

EP-A1- 0 154 539 WO-A-87/02671

EP-A- 0 171 496

EP-A- 0 184 187

- · Journal of immunology (Baltimore) Volume 137 Number 3, issued 01 August 1986, SAHAGAN et al. "A genetically engineered murine human chimeric antibody retains specificity for human tumor-associated antigen See pages 1066-1074.
- Nature (London) Volume 309, 24 May 1984, SHARON et al. "Expression or VHCK chemaeric protein in mouse myeloma cells" See pages 364-367.
- Nature(London) Volume 312, issued 13 December 1984. Boullanne et al. "Production of functional chimaeric mouse/human antibody* see pages 643-646.
- Proceedings of the National Academy of Science USA (Washington, D.C.) Volume 81, Issued June 1984, Cabiliv et al. "Generation of antibody activity from immunoglobulin chains produced in Escherichia coli" see page 3273-3277.
- Biotechniques (Natick, MA) Volume 4 issued May-June 1986, Oi et al "Chimeric Antibodies" see pages 216-220.
- · Nature (London) Volume 314, Issued 04 April 1985, TAKEDA et al. "Construction of chimaeric processed immunoglobulin genes containing mouse variable and human constant region sequences" see pages 452-454
- Proceedings of the National Academy of Sciences, USA (Washington, D.C.) Volume 81 Issued November 1984, MORRISON et al., *Chimeric human antibody molecules : mouse antigent-binding domains with human constant region domains" see pages 6851-6855.
- · Science, (Washington, DC) Volume 229 Issued 20 September 1985, MORRISON *Transfectomas provide novel chimeric antibodies" see pages 1205-1207.

(Cont. next page)

- Science, (Washington, D.C.) Volume 240 issued 20 May 1988, BETTER et al. pages 1041-1043
- Nucleic Acids Research, (Oxford) Volume 12 Issued December 1984, KRAMER et al. *The gapped duplex DNA approach to ollgonucleotide-directed mutation
- construction". See pages 9441-9455.

 Bio/Technology (Martinsville, NJ) Volume 4 issued November 1986, HSIUNG et al.
- "High-level expression efficient secretion and folding or human growth hormone in Escherichia coil" see pages 991-995 particularly p. 992 and floure 3.
- Nature (London) Volume 312 Issued 13 December 1984, NEUBERGER et al. "Recombinant antibodies possessing novel effector functions" see pages 604-608,
- particularly pages 605-606 and figures 1 and 2.

 Nature (London) Volume 314 Issued 21 March 1985, NEUBERGER et al. "A hapten-specific chimaeric igE antibody with human physiological effector function" see pages

268-270.

 Journal of Immunology (Baltimore) Volume 135 Number 5 Issued November 1985, TAN et al. "A human-mouse chimeric immunologicbulin gene with a human variable region is expressed in mouse mysloma cells" see pages 3564-8567.

- Gene, (Amsterdam) Volume 43. Issued 25
 August 1986, Williams et al. "Production of antibody-tagged enzymes by myeloma cells: application to DNA po-lymerase I Klenow fragment" see pages 319-324.
- Nature, (London) Volume 314 o4 April 1985, Wood et al. "The synthesis and in vivo assembly of functional antibodies in yeast" see pages 446-449.
- Proc. Natl. Sci USA 80 (1983), pp 6351-6355.
- Proc. Nati. Acad. Sci. USA 84 (1987), pp. 3439-3443.
- The EMBO Journal, 44, No. 1, pp. 519-526, 1985;
 McCarthy et al
- Amino Acid and Nucleotide Biosynthesis, pp. 284-287; Jones E.W.
- PNAS, 83, pp. 8506-8510; Schoner et al
- Methods in Enzymology, Goeddel, ed. Academic
- Press, 1990, pp. 166-187; Stader et al.
 Protein Engineering, 2(3), 1988, pp. 169-170;
- Blo/Techn., 10, 1992, pp. 163-167; Carter et al
- Nucl. Acid Res., 12, 1984, pp.3791-3806; Boss et
- Science, 240, 1988, pp. 1038-1041; Skerra et al
- PNAS, 52, 1964, pp. 1099-1166; Haber et al
 Adv. Immunol., 44, 1989, pp. 65-92; Morrison & Ol

Remarks:

Divisional application 93100041.8 filed on 25/07/88

Description

Field of the Invention

[0001] This invention relates to recombinant DNA methods of preparing immunoglobulins, genetic sequences coding therefor, as well as methods of obtaining such sequences.

Background Art

- [0002] The application of cell-to-cell fusion for the production of monoclonal antibodies by Kohler and Milistani, (Nature (London), 286: 495, 1975) has paymed a revolution in biology equal in impact to the invention of recombinant DNA cloning. Hybridoma-produced monoclonal antibodies are almady widely used in clinical diagnoses and basic scientific studies. Applications of human B cell hybridoma-produced monoclonal antibodies hold great promise for the clinical treatment of cancer, viral and microbial infections, B cell immunodeliciencies with climinished antibody production, and other diseases and disorders of the immune system.
- [0003] Unfortunately, yields of monoclonal antibodies from human hybridoma cell lines are relatively low (1 ug/ml in human x human compared to 100 ug/ml in mouse hybridomas), and production costs are high for antibodies made in large scale human issue culture. Mouse x muse hybridomas, on the other hand, are useful because they produce abundant amounts of protein, and these cell lines are more stable than the human lines. However, repeated injections
- of Toreign' antibodies, such as a mouse antibody, in humans, can lead to harmful hypersensitivity reactions. [0004] There has therefore been recent exploration of the possibility of producing antibodies having the advantages of monoclonals from mouse-mouse hybridomas, yet the species specific properties of human monocional antibodies. [0005] Another problem faced by immunologists is that most human monocional antibodies (i.e., antibodies having human recognition properties) obtained in cell culture are of the light type. When it is desirable to obtain human monocionals of the light type, however, it has been necessary to use such techniques as cell sorting, to separate the few cells which have switched to producing antibodies of the light type. A need therefore exists for a more ready method of switching antibody classes, for any given antibody of a predetermined or desired antigenic specificity.
- [0006] The present invention bridges both the hybridoma and monoclonal antibody technologies and provides a quick and efficient method, as well as products derived therefrom, for the improved production of chimeric human/monhuman antibodies, or of "class switched" antibodies.
 - [0007] Approaches to the problem of producing chimeric antibodies have been published by various authors.
- [0008] Morrison, S. L. et al., Proc. Natl. Aced. Sci., USA, 81: 6851-6855 (November 1984), describe the production of a mouse-human antibody molecule of defined antigen brinding specificity, produced by joining the variable region of a mouse antibody-producing myeloma cell line with known antigen brinding specificity to human immunoglobulin constant region genes using recombinant DNA techniques. Chimeric genes were constructed, wherein the heavy chain variable region exon from the myeloma cell line S107 well joined to human (gG1 or [gG2 heavy chain constant region exons, and the light thain variable region exon from the myeloma cell lines 100 well joined to human (gG1 or [gG2 heavy chain constant region exon, and the light thain variable region exon from the same myeloma to the human kappa light chain exon.

 These genes were transfected into mouse myeloma cell lines and transformed cells producing chimeric mouse-human antiphosphorcholine antibodies were thus developed.
- [0009] Morrison, S. L. <u>et al.</u>, European Patent Publication No. 173494 (published March 5, 1986), disclose chimeric 'receptors' (e.g. antibodies) having variable regions derived from one species and constant regions derived from another. Mention is made of utilizing cDNA cloning to construct the genes, although no details of cDNA cloning or priming are shown. (see pp 5, 7 and 6).
- 45 [0010] Boullanne, G. L. et al., Nature, 312: 643 (December 13, 1984), also produced antibodies consisting of mouse variable regions joined to human constant regions. They constructed immunoglobulin genes in which the DNA segments encoding mouse variable regions specific for the hapten trintiropheny/ (TNP) were joined to segments encoding human <u>mu</u> and <u>kappa</u> constant regions. These chimeric genes were expressed as functional TNP binding chimeric IgM. [0011] For a commentary on the work of Boullanne <u>et al.</u>, and Morrison <u>et al.</u>, ase Murro, <u>Nature</u>, 312: 597 (December
- 0 13, 1984), Dickson, <u>Genetic Engineering News</u>, 5, Mo. 3 (March 1985), or Marx, <u>Science</u>, <u>229</u>, 455 (August 1985), [0012] Neuberger, M. S. <u>et al.</u>, <u>Nature</u>, <u>311</u>: 268 (March 25, 1986), also constructed a chimeric heavy chain immunoglobulin gene in which a DNA segment encoding a mouse variable region specific for the hapten 4-hydroxy-3-nitrophenacety (NP) was pioned to a segment encoding the human <u>engilor</u> region. When this chimeric gene was transfected into the <u>1955</u>, cell line, an antibody was produced which bound to the NP hapten and had human LGE properties.
- 55 [0013] Neuberger, M.S. gt al., have also published work showing the preparation of cell lines that scerete haptenspecific antibodies in which the Fc portion has been replaced either with an active enzyme moiety (Williams, G. and Neuberger, M.S. <u>Gene 43</u>:319, 1986) or with a polypeptide displaying c-myc antigenic determinants (<u>Nature</u>, <u>312</u>:804, 1984).

- [0014] Neuberger, M. et al., PCT publication WO 86/01533, (published March 13, 1986) also disclose production of chimeric antibodies (see p. 5) and suggests, among the technique's many uses the concept of "class switching" (see p. 6).
- [0015] Taniguchi, M., in European Patent publication No. 171 496 (published February 19, 1995) discloses the production of chimeric antibodies having variable regions with tumor specificly derived from experimental animals, and constant regions derived from human. The corresponding heavy and light chain genes are produced in the genomic form, and quoressed in mammalian calls.
- [0018] Takeda, S. gt <u>al.</u>, Nature, 314. 452 (April 4, 1985) have described a potential method for the construction of chimeric immunoglobulin genes which have intron sequences removed by the use of a retrovirus vector. However, an our unexpected spike donor site caused the deletion of the V region leader sequence. Thus, this approach did not yield complete chimeric antibody molecules.
 - (0017) Cabilly, S. <u>et al.</u>, <u>Proc. Natl. Acad. Sci., USA, 81</u>: 3273-3277 (June 1984), describe plasmids that direct the synthesis in <u>E. coli</u> of heavy chains and/or light chains of anti-carcinoembryonic antigen (CEA) antibody. Another plasmid was constructed for expression of a truncated form of heavy chain (Fd) frament in E. coli Functional CEA-binding divides constructed for expression of a truncated form of heavy chain (Fd) frament in E. coli Functional CEA-binding divides constructed for expression of a truncated form of heavy chain (Fd) frament in E. coli Functional CEA-binding divides the form of the procession of the form of
- is activity was obtained by in vitro reconstitution, in E. coll extracts, of a portion of the heavy chain with light chain. [0018] Cabilly, S. gt all, European Pattern Publication 15203 (published November 14, 1984) describes chimeric immunoglobulin genes and their presumptive products as well as other modified forms. On pages 21, 28 and 33 it discusses cDNA chonina and primina.
- discusses cDNA cloning and priming.

 [0019] Boss, M. A., European Patent Application 120694 (published October 3, 1984) shows expression in <u>E. coli</u>
 of non-chimeric immunoglobulin chains with 4-nitrophenyl specificity. There is a broad description of chimeric antibodies
 - but no datalis (see p. 9).

 [0020] Wood, C. R. gl. <u>M. Nature</u>, 314: 446 (April, 1985) describe plasmide that direct the synthesis of mouse anti-NP antibody proteins in yeast. Heavy chain <u>mu</u> antibody proteins appeared to be glycosylated in the yeast cells. When both heavy and light chains were synthesized of in the same cell, some of the protein was assembled into functional
- antibody molecules, as detected by anti-NP binding activity in soluble protein prepared from yeast cells, [0021] Alexander, A. et al., Proc. Nat. Acad. Sci. USA, 79: 3260-3264 (1982), describe the preparation of a cDNA sequence coding for an abnormally short human in gamma heavy chain (DMM gamma 3 HCD serum protein) containing a 19- amino acid leader followed by the first 15 residues of the V region. An extensive internal deletion removes the remainder of the V and the entire C₄1 domain. This is cDNA coding for an internally deleted molecule.

- 50 [0022] Doby, T. W. et al., Proc., Natl. Acad. Sci., USA, 27: 6027-6031 (1980), describe the preparation of a cDNA sequence and recombinant plasmids containing the same coding for mu and <u>Asopa</u> human immunoglobulin polypepides. One of the recombinant DNA molecules contained codons for part of the CH₃ constant region domain and the entil of noncodina sequence.
- [0023] Seno, M. at al., Nucloic Acids Research, 11: 719-726 (1983), describe the preparation of a cDNA sequence 35 and recombinant plasmids containing the same coding for part of the variable region and all of the constant region of the human IgE heavy chain (peption chain).
 - [0024] Kurokawa, T. et al., |bid, 11: 3077-3085 (1983), show the construction, using cDNA, of three expression plasmids coding for the constant portion of the human IgE heavy chain.
- [0025] Liu, F. T. et al., Proc. Nat. Acad. Sci., USA, 81: 5369-5373 (September 1984), describe the preparation of a cDNA sequence and recombinant plasmids containing the same encoding about two-thirds of the CH₂, and all of the C_N3 and C_N4 domains of human (gE heavy chain.
 - [0026] Tsujimoto, Y. et al., Nucleic Acids Res., 12: 8407-8414 (November 1984), describe the preparation of a human V lambda cDNA sequence from an Ig lambda-producing human Burkitt lymphoma cell line, by taking advantage of a cloned constant region gene as a primer for CDNA synthesis.
- 45 [0027] Murphy, J., PCT Publication WO 83/03971 (published November 24, 1983) discloses hybrid proteins made of fragments comprising a toxin and a cell-specific ligand (which is suggested as possibly being an antibody).
 - [0028] Tan, et al., J. Immunol. 135:8564 (November, 1985), obtained expression of a chimeric human-mouse Immunoglobulin genomic gene after transfection into mouse myeloma cells.
- [0029] Jones, P. T., et al., <u>Nature 321</u>:552 (May 1986) constructed and expressed a genomic construct where CDR domains of variable regions from a mouse monoclonal antibody were used to substitute for the corresponding domains in a human antibody.
 - [0030] Sun, L.K., at al., <u>Hybridoma 5 suppl.</u> 1517 (1986), describes a chimeric human/mouse antibody with potential tumor specificty. The chimeric heavy and light chain genes are genomic constructs and expressed in mammalian cells. [0031] Sahagan <u>et al.</u>, <u>J. Immum.</u> 137:1066-1074 (August 1986) describe a chimeric antibody with specificity to a
- Wman turnor associated antigen, the genes for which are assembled from genomic sequences. [0032] For a recent review of the field see also Morrison, S.L., <u>Science</u> 229: 1202-1207 (September 20, 1985) and Oi, V.T., et al., BioTechniques 4:214 (1986).
 - [0033] The Oi, et al., paper is relevant as it argues that the production of chimeric antibodies from cDNA constructs

in yeast and/or bacteria is not necessarily advantageous.

[0034] See also Commentary on page 835 in Biotechnology 4 (1986).

SUMMARY OF THE INVENTION

5

25

35

40

45

60

[0035] The invention provides a novel approach for producing genetically engineered antibodies of desired variable region specificity and constant region properties through gene cloning and expression of light and heavy chains. The cloned immunoglobulin gene products can be produced by expression in genetically engineered organisms.

[0036] The application of chemical gene syntheses, recombinant DNA cloning, and production of specific immunopbullin chains in various organisms provides an effective solution for the efficient large scale production of human monoclonal antibodies. The invention also provides a solution to the problem of class switching antibody molecules, so as to readful versear immunoplobulins of a contain briddin seperificity of any other class.

[0037] According to the present invention there are provided polynucleotide molecules comprising a prokayotic promoter region in operable inkage to a dicistronic transcription unit, the unit necoding a heavy chain immunoglobulin, or a fragment thered, and a light chain immunoglobulin, the heavy and light chains being separately operably linked to a sequence coding for a polyneptic secretion signal.

[0038] The invention also relates to such recombinant molecules in vehicles such as plasmid vectors, capable of expression in desired prokaryotic hosts.

[0039] The invention provides prokaryotic hosts capable of producing, by culture, chimeric antibodies and methods of using these hosts

[0040] The invention may be used to provide chimeric immunoglobulin individual chains, whole assembled molecules, and immunoglobulin fragments (such as Fab) having human constant regions and non-human variable regions, wherein both variable regions have the same bindin ospecificity.

[0041] Among other immunoglobulin chains and/or molecules that can be provided by the invention are:

- (a) a complete functional, immunoglobulin molecule comprising:
 - (i) two identical chimeric heavy chains comprising a non-human variable region and human constant region and
 - (ii) two identical all (i.e. non-chimeric) human light chains.
- (b) a complete, functional, immunoglobulin molecule comprising:
 - (i) two identical chimeric heavy chains comprising a non-human variable region and a human constant region, and
 - (ii) two identical all (i.e. non-chimeric) non-human light chains.
- (c) a monovalent antibody, i.e., a complete, functional immunoglobulin molecule comprising;
 - (i) two identical chimeric heavy chains comprising a non-human variable region and a human constant region, and
 - (ii) two different light chains, only one of which has the same specificity as the variable region of the heavy chains. The resulting antibody molecule binds only to one end thereof and is therefore incapable of divalent binding;
- (d) an antibody with two different specificities, i.e., a complete, functional immunoglobulin molecule comprising:
 - (f) two different chimeric heavy chains, the first one of which comprises a non-human variable region and a human constant region and the second comprises a different non-human variable region, and a human constant region, and
 - (ii) too different chimate light chains, the first one of which comprises a non-human variable region having the same specificity as the first heavy chain variable region, and a human constant region, and the second comprises a non-human variable region having the same specificity as the second heavy chain variable region, and a human constant region.
- 55 [0042] The resulting antibody molecule binds to two different antigens.
 - [0043] The invention also provides for the production of functionally active chimeric immunoglobulin fragments secreted by prokaryotic or eukaryotic hosts or fully folded and reassembled chimeric immunoglobulin chains.
 - [0044] Genetic sequences, especially cDNA sequences, coding for the aforementioned combinations of chimeric

chains or of non-chimeric chains are also provided herein.

[0045] The use of cDNA sequences is particularly advantageous over genomic sequences (which contain introns), in that cDNA sequences can be expressed in bacteria or other hosts which lack RNA splicing systems.

[0046] Among preferred specific antibodies are those having specificities to cancer-related antigens.

BRIEF DESCRIPTION OF THE FIGURES

5

[0047] FIGURE 1 shows the DNA rearrangements and the expression of immunogboulin <u>mu</u> and <u>gamma</u> heavy chain genes. This is a schematic representation of the human heavy chain gene complex, not shown to scale. Heavy chain variable V region formation occurs through the joining of V_{In}. D and J_{In} gene segments. This generates an active <u>mu</u> gene. A different kind of DNA rearrangement called 'class switching' relocates the joined V_{In}. D and J_{In} region from the <u>mu</u> constant C region to another heavy chain C region (switching to <u>gamma</u>) is diagrammed here). The scheme empaisizes that the J region is a common feature of axis consequence of the common feature of expressed lines. The J region is also a common feature of expressed lines are common entered to the common feature of expressed lines are common entered to the common feature of expressed lines are common entered to the consequence of the common feature of expressed lines are common feature of expressed lines are common entered to the common feature of expressed lines are common entered to the common feature of expressed lines are common entered to the common feature of expressed lines are common entered to the common feature of expressed lines are common entered to the common feature of expressed lines are common entered to the common feature of expressed lines are common entered to the common feature of expressed lines are common entered to the common feature of expressed lines are common entered to the commo

5 [0048] FIGURE 2 shows the known nucleotide sequences of human and mouse J regions. Consensus sequences for the J regions are shown below the actual sequences. The oligonucleotide sequence below the mouse <u>leapse</u> J region consensus sequence is a Universal Immunoglobulin Gene (UIG) oligonucleotide which is used in the present invention.

[0049] FIGURE 3 shows a scheme noting the use of the UIG oligonucleotide primer for the synthesis of cDNA complementary to the variable region of immunoglobulin messenger RNA, or the use of oligo-dT as a primer for cDNA synthesis, followed by in vitro mutagenesis.

[0050] FIGURE 4 shows the synthesis and analysis of human IgG1 genes, including three Isolated clones (A.b), one of which (pGMH-6) is utilized as a cloning vector (B). A 1.5 kb deletion of pBR322 sequence between <u>Barn</u> HI and Pyull is marked. Not to scale.

25 [0051] FIGURE 5 shows the cloning vector pQ23, a modified pBR322, useful for cDNA cloning at the KpnI site. This vector also contains the useful restriction enzyme sites BallI plus Sall. Not to scale.

[0052] FIGURE 6 shows in A. the synthesis and analysis of human light chain <u>kappa</u> genes. The Figure also shows in B. (not in scale) construction of a human C_x region cloning vector pING2001.

[0053] FIGURE 7 shows primers designed for immunoglobulin V region synthesis. (A) shows the heavy chain J-C regions and primers. A DNA version of each mouse J heavy region is shown directly above primers designed from that sequence. Mouse J regions are 5' to 3', left to right, while primers are 3' to 5', left to right. Primer names are included in brackets, and numbers of mucleotides (N) and number of misenabethes with each J_A; region are islead to the right. Primers which introduce a <u>Bul</u>EI site are underlined. (3) shows the light chain J regions and primers. The same as for (A) except for light chains. Primers designed to introduce a <u>Bul</u>EI site are underlined, as is the <u>Bul</u>EI site present in 3P INC2016E. (C) shows mouse variable region consensus UIG primers. The actual primer sequence is shown below that consensus sequence. The human C_K <u>Hin</u>dIII vector pGML60 is shown below. (D) shows a mouse gamma 2a J/C junction primer.

[0054] FIGURE 8 shows the synthesis of heavy chain V region module genes using oligonucleotide primed cDNA synthesis. Not to scale.

(0055) FIGURE 9 shows the construction of hybrid mouse-human immunoglobulin genes. Panel A shows construction of a heavy chain gene. Stippled regions show C region modules, while hatched or black regions show V region modules. Not to scale.

[0056] FIGURE 10 shows the construction of cDNA cloning-expression shuttle vectors for mammalian cells. The vectors pilot(2003 and pilot(20038 and pilot(20038) and pilot(20038) and pilot(20038) pilot(

[0057] FIGURE 11 shows the construction of the heavy chain expression plasmid plNG2006E. Arrows show SV-40 promoter locations and directions of transcription. Hatched and black areas show mouse V region modules, while stippled areas show human C region modules. Not to scale.

[0058] FIGURE 12 shows the structure of the chimeric anti-hapatitis heavy chain genes in the expression plasmids pin/32008E and pin/32012E. Panel A shows the structure of mouse human chimeric anti-hapatitis heavy chain genes pin/32008E and pin/32012E. Panel A shows the structure of mouse human heavy chain constant region cDNA clone pGMH-8 and the mouse heavy chain variable region cDNA chones pBS13-1 and pJ8-11 were used to make the hybrid gene used in pin/32008E. Halched gene blocks indicate mouse variable region sequences, while open gene blocks show human IgG1 constant region sequences. Panel B shows the nucleotide sequence of the anti-hapatitis B heavy chain variable region capital profices and pin/32008E. Halched post blocks are profit by the sequence of the anti-hapatitis B heavy than variable region in pin/32006E and pin/32012E. pin/32012E was constructed by first insering a Stull steaty

Sall six of pING1322 (See Figure 16) to form pING13228gll. The chimeric heavy chain gene from this plasmid was inserted into the expression vector pING2009E if, resulting in pING2012E, pING2012E differs from pING 2008E into region immediately upstream of the initiator ATG. Underlined nucleotides denote human J region sequences from the CONA clone pCNM+5, afterisked amino acid 117 indicates as ingle change at this site from mouse to human sequence (Ala to Ser) introduced in the chimeric grain of the condition of

[0059] FIGURE 13 shows in panel A the J-C junction region nucleotide sequence in light chain clones derived from plNc2001 (pMACK-3, plNc2013E, plNc2007E, plNc2010E-gpt and plNc2014E-gpt). The J region sequence originating from pK2-3 is marked human JK4. The 6 nucleotide not predicted by genomic sequencing is marked with an asterisk. The oligonucleotide primer (K2-4BCLI) used to modify this sequence is shown below the human JK4 se-

quence. Panel B diagrams the method of site-directed multagenesis used to make pING2016E-gpt. Not to scale. [0060] FIGURE 14 Gene copy number of the transfected sequences in two transformants. nDNA from 2AE9, 2BH10 were digested with the enzymes indicated. The concentration of DNA is titrated down across the lanes with the amount indicated above them. The probe contains human C garmma 1 sequences (print/e24 Aget-BamHI). The reference is germ-line or 6MM2146 nDNA digested with Aget. The 3" Agai site is 2 by beyond the site of poly(A) addition (3).

[0061] FIGURE 15 shows the nucleoticle sequence of the V region of the LS V_h cDNA clone pH3-8a. The sequence was determined by the diffeorytemination method using M13 subctones of gene fragments (shown below). Open circles denote armino acid residues confirmed by peptide sequence. A sequence homologous to D₈₊₂ in the CDR3 region is underlined.

(9062) FIGURE 16 shows the nucleotide sequence of the V region of the L6 V_K cDNA clone pt.3-12a. The oligonucleotide primer used for site-directed multagenesis is shown below the J_K5 segment. Open circles denote amino acid residues confirmed by peptide sequence.

[063]. FIGURE 17 shows the construction of chimeric L6V₄, plus human C gamma expression plasmids. Panel (a) shows the sequences of the BAL-31 deletion clones M13mp19-C1-delta 4 (C1-delta 4) and M13mp19-C1-delta 2 (C1-delta 2). The 5° and of the cDNA clone, pH-5-6a, is denoted with an arrow. M13 sequences are underlined. The oligonucleotide primer used for this experiment is H3-6a (5° - GACTGCACATGGG-3), which primes in F1 near the nature N terminus. Panel (b) shows the strategy for site-directed mutagenesis of 1 ug of clones C1-delta 4 and C1-delta 21, each annealed to 20 ng of the 31-mer oligonucleotide MUH2-Apal. Complementary strand synthesis with the Klenow fragment of L0NA polymerase was ar troom temperature for 30 mil, then 11°C for 27 hours. Transfected phage plaques were adsorbed to nitrocallulose, fixed with NaOH, and hybridized to ³²P-abelted MUH2-Apal oligonucleotide at 65°C, 18 hours, in 47158 (0.6 M NaC1, 0.0 4M TF1A+C) (pH 7.4), 0.00 4M EDTA) plus 10°C dextran sulfate. Final was of the filters was at 65°C, 4x5SPE, 0.1% SDS for 15 min. (Maniatis, T, gt al, Molecular Cloning: A Laboratory Manual 1982). Positive plaques were detected by overnight exposure to Kodak XAR filtin, and were directly plicked for growth and restriction enzyme analysis of FF DNA. Mismatches of the MIH2-Apal oligonucleotide to the mouse C_{p1} are denoted. Sea the changes shown below the oligonucleotide. Panel (c) shows the strategy of the substitute of the MIH2-Apal oligonucleotide to the mouse C_{p1} are denoted. Sea the clones are constanted to the constanted of the first of the MIH2-Apal oligonucleotide to the mouse C_{p1} are denoted. Sea the clones are constanted to the constanted of the clones are constanted to the constanted of the con

of each of the mutagenized L6-V_H modules for the resident V_H of the chimeric expression plasmid pING2012 to generate pING2111 and pING2112.

[0064] Fig.UNE 18 shows the construction of the chimeric L6 expression plasmid pING2119. The <u>Sall</u> to <u>Bam</u>HI

fragment from pING2100 is identical to the <u>Sall</u> to <u>Bam</u>HI A fragment from pING2012E.

[0065] FIGURE 19 shows the modification of the V_K gene and its use in constructing light chain and heavy plus light chain aversation plasmids.

45

(a) Deletion of the oligo d[GC] segment 5' of V_K of L6. The oligonuclectide is a 22-mer and contains a <u>Sall</u> site. The 3 mismatches are shown. The V_K gone, after mutagenesis, is joined as a <u>Sall-Hind</u>III fragment to the human C K module. The expression plasmid thus formed is piNG2119.

(b) pING2114, a heavy plus light chain expression plasmid. The expression plasmid pING2114 contains the L6 heavy chain chimeric gene from pING2111 and the chimeric light chain from pING2119 (bold line).

[0056] FIGURE 20 shows a summary of the sequence alterations made in the construction of the L6 chimeric antibody expression plasmids. Residues underlined in the 5' untranslated region are derived from the cloned mouse kapps and heavy-chain genes. Residues circled in the VIC boundary result from mutagenesis operations to engineer restriction enzyme sites in this region. Residues denoted by small circles above them in the L6 heavy-chain chimera also result from mutagenesis. They are gient changes.

[0067] FIGURE 21 shows the 2H7 V_H sequence. The V_H gene contains J_H1 sequences and DSP.2 sequence elements. Small circles above the amino acid residues are those that matched to peptide sequences.

[0068] FIGURE 22 shows the 2H7 V_L sequence. The V_K gene contains J_L 5 sequences. A 22-mer oligonucleotide was used to place a Sall site 5' of the ATG initiator codon. Small circles above the amino acid residues are those that matched to peptide sequences.

[0069] FIGURE 23 shows the chimeric immunoglobulin gene expression plasmids of the 2H7 specificity. One gene plasmids are piNG2101 (V_{th}.neo), piNG2106 (V_{th}.neo) and piNG2107 (V_{th}.gpt). The others are two-gene plasmids. Their construction involved the ligation of the targer <u>Ndel</u> tragments of piNG2107 and piNG2107 to linearized piNG2106 partially digested with <u>Ndel</u>, pHL2-11 and pHL2-26 were obtained from piNG2107 and piNG2105 and piNG2106 and piNG2106 to the ndel phone of the phone of the ndel phone of the ndel phone of the phone of the

[0070] FIGURE 24 shows a summary of the nucleotide changes introduced in the V_R and V_K in the construction of the chimeric plasmids. The cognate V_R and V_K nucleotide residues in the 5' end are underlined. Circles residues in the V_R unclose are derived from the human C modules.

[0071] FIGURE 25 shows the strategy used to fuse the mature L6 chimeric light chain sequence to the yeast invertase signal sequence and shortened PGK promoter. The open clothel line represents yeast invertase signal sequence DNA. The solid double line represents yeast PGK DNA; > represents the PGK promoter, -I represents the PGK terminator; RF = Replicative Form, piNG 1225 was derived by fusing human C₆ DNA to the PGK promoter, piNG 1149 was derived by fusing the yeast invertace signal sequence to the yeast PGK promoter. (A) shows the strategy for introduction by mitter omutageness of an Agtill site in the signal sequence processing site. (B) shows the DNA sequence of the single-stranded mutagenesis primer and the corresponding numbagenized DNA sequence. (C) shows the strategy used to construct a plasmid containing the mature light chain sequence fused to the invertase signal sequence and shortened PGK promoter.

[0072] FIGURE 26 shows the strategy used to fuse the mature L5 chimeric heavy chain sequence to the yeast invertase signal sequence and shortened PGK promoter, pIMG1298 contains the chimeric heavy chain gene with the variable region from the 2H7 mouse monoclonal antibody (see example IV). All symbols are as defined in legend for Figure 25. (A) shows the strategy for introduction by <u>in vito</u> mutagenesis of an <u>Sall</u> site in the signal sequence processing site. (8) shows the DNA sequence of the single-stranded mutagenesis primer and the corresponding unmutagenized DNA sequence. (C) shows the strategy used to construct a plasmid containing the mature heavy chain sequence fused to the invertage single signal sequence and shortened PGK promoter.

65 [0073] FIGURE 27 shows the strategy used to remove non-yeas! 3' untranslated DNA sequences from the L6 chimaric light chain gene and to construct a plasmid containing the light chain gene fused to the invertase signal sequence and shortened PGK promoter in which all sequences are either known by DNA sequence analysis or proven to be functional. pBR322NA is derived from pBR322 by deletion of DNA from Ndel to Aual. Symbols are as defined in legend for Figure 25.

20 [0074] FIGURE 28 shows the strategy used to remove non-yeast 3' untranslated DNA sequence from the L6 chimeric heavy chain gene and to construct a plasmid containing the heavy chain gene fused to the invertase signal sequence and shortened PGK promoter in which all sequences are either known by DNA sequence analysis or proven to be functional. Symbols are as defined in legend for Figure 25.

[0075] FIGURE 29 shows the strategy used to clone the L6 chimeric light chain gene fused to the invertase signal sequence and shortened PGK promoter into yeast-<u>E, col</u>l shuttle vectors containing the PGK transcription terminationpolyadenylation signal, yeast replication sequences, and genes for selection of transformants. Symbols are as defined in legend for Figure 25.

[0076] FIGURE 30 shows the strategy used to clone the L5 chimeric heavy chain gene fused to the invertase signal sequence and shortened PoK promoter into yeast—E_col shuttle vectors containing the PSK transcription terminationpolyadenylation signal, yeast replication sequences, and genes for selection of transformants. Symbols are as defined in legend for Figure 25.

[0077] FIGURE 31 shows a schematic diagram of the structure of human IgG1.

[0076] FIGURE 32(A) shows the strategy used to htroduce a stop codon and Bell site into the hinge region of human gamma 1. (B) shows the DNA sequence of the single-stranded primer used for <u>in vitre</u> multagenesis of the gamma-1 hinge region and the corresponding unmutagenized sequence. Vertical arrows represent inter-chain disulfide bonds. Symbols are as defined in legend for Figure 25.

[0079] FIGURE 33 shows the strategy used to fuse the L6 chimeric heavy chain gene containing a stop codon in the hinge region (Fd dhain) to the yeast invertase signal sequence and shortened PGK promoter. Symbols are as defined in legend for Figure 25.

0 [0080] FIGURE 34 shows the strategy used to remove nonyeast 3' untranslated sequences from the L6 chimeric Fd chain and to construct a plasmid containing the Fd chain fused to the invertises signal sequence and shortened PGK promoter in which all sequences are either known by DNA sequence analysis or proven to be functional. Symbols are as defined in legend for Figure 25.

[0081] FIGURE 25 shows the strategy used to clone the L6 chimeric Fd chain gene fused to the invortace signal sequence and shortened PGK promoter into yeast—E_colis buttle vectors containing the PGK transcription termination-polyadenylation signal, yeast replication sequence, and genes for selection of transformants. Symbols are as defined in legend for Ficure 25.

[0082] FIGURE 36(A) shows the nucleotide sequence surrounding the N-terminus of the Erwinia caratovora pelB

gene (Lei, S.P., et al., J. Baderiol, (1987, in press)). The Ndel and Haell I sites used in cloning are shown. The arrow indicates the leader poptidase cleavage site for poctate lyase. (8) shows the cloning strategy for construction of the pelB leader cartridge, pSS1004 contains a 1.9 kb <u>Dral</u> fragment cloned into the <u>Smal</u> site of pUCB. Symbols are defined in the leagend for Figure 39.

5 [0083] FIGURE 37 shows the construction of light chain expression plasmids pRR177-8, pRR180, pRR190, and pRR191. In addition to the plasmids described in the total, MT3mpl B and pIT 181 were used pT181 contains the mature light chain gene fused directly following the ATS initiation codon of the <u>art80</u> gene in pT2 (see Figure 40). [0084] FIGURE 38 shows the construction of Fet expression desards pRI747-95. pPR180, and pRR195.

[0085] FIGURE 39 shows the restriction maps of the light chain and Fd gene cassette in pFK100, pFK101, pFK102, o pFK103, and pFK104. These plasmids were constructed as described in the text using plasmids outlined in Figure 37 and 38. The arrow indicates the direction of transcription from the <u>lag</u> promoter. <u>FL caratevova</u> and eukanyotic noncoding sequences are shown as open bars. The <u>pelB</u> leader sequence is cross-hatched and the closed bar represents the antibody genes Ed and flight chain (K).

[0085] FIGUFE 40(A) shows the construction of a vector for arabinose inducible Fab expression. Plasmid pTI2 (Masson and Ray, Nucl. Acids Res., 14:5693 (1986)) is a 6431 bp plasmid encoding the araC gene, the araB percentor and a portion of the araB gene from pNG1 (Johnston, S., et al., Gene 34:134 (1985)) in a derivative of pBR322. An Nccl site has been engineered at the ATG initiation codon of the araB gene. (B) shows the introduction of the laci gene into pFK102.

20 DESCRIPTION OF THE PREFERRED EMBODIMENTS

INTRODUCTION

[0087] Generally, antibodies are composed of two light and two heavy chain molecules. Light and heavy chains are divided into domains of structural and functional homology. The variable regions of both light (V_c) and heavy (V_s) chains determine recognition and specificity. The constant region domains of light (C_j) and heavy (C_s) chains confer important biological properties such as antibody chain association, secretion, transplacental mobility, complement binding, and the like.

[0088] A complex series of events leads to immunocjobulin gene expression in B cells. The V region gene sequences or conterring antigen specificity and binding are located in separate germ line gene segments called V_H. D and J_H, or V_L and J_L. These gene segments are joined by DNA rearrangements to form the complete V regions expressed in heavy and light chains respectively (Figure 1). The rearranged, joined (V_L-J_L and V_H-D-J_H)V segments then encode the complete variable regions or anticen binding domains of light and heavy chains, respectively.

35 DEFINITIONS

50

[0089] Certain terms and phrases are used throughout the specification and claims. The following definitions are provided for purposes of clarity and consistency.

- 40 1. Expression vector a plasmid DNA containing necessary regulatory signals for the synthesis of mRNA derived from gene sequences, which can be inserted into the vector.
 - 2. Module vector a plasmid DNA containing a constant or variable region gene module.
 - 3. Expression plasmid an expression vector that contains an inserted gene, such as a chimeric immunoglobulin gene.
 - 4. Gene cloning synthesis of a gene, insertion into DNA vectors, and identification by hybridization and the like.

 5. Transfection the transfer of DNA into mammalian cells.
 - Promoter region a nucleotide sequence which provides a cell with the regulatory sequences needed to express
 - an operably linked cistron or operon.
 - Secretion signal a polypeptide present at the N-terminus of a chimeric immunoglobulin chain useful in aiding in the secretion of the chain to the outside of the host. Also called "leading peptide." or "leader."

GENETIC PROCESSES AND PRODUCTS

[0090] The invention provides a novel approach for the cloning and production of human antibodies with desired specificity, Generally, the method combines five elements:

(1) Isolation of messenger RNA (mRNA) from B cell hybridoma lines producing monoclonal antibodies against specific antigens, cloning and cDNA production therefrom;

- (2) Preparation of Universal Immunoglobulin Gene (UIG) oligonucleotides, useful as primers and/or probes for cloning of the variable region gene segments in the light and heavy chain mRNA from specific human or nonhuman hybridoma cell lines, and cDNA producton therefrom.
- (3) Preparation of constant region gene segment modules by cDNA preparation and cloning, or genomic gene preparation and cloning.
- (4) Construction of complete heavy or light chain coding sequences by linkage of the cloned specific immunoglobulin variable region gene segments of part (2) above to cloned human constant region gene segment modules;
- ulin variable region gene segments of part (2) above to cloned numan constant region gene segment modules, (5) Expression and production of light and heavy chains in selected prokaryotic hosts, either in separate fermentations followed by assembly of antibody molecules in <u>vitro</u>, or through production of both chains in the same cell.
- [0091] The invention employs cloned hybridoma B cell lines producing monoclonal antibodies of defined specificity for the isolation of mRNA for cDNA cloning. Because many lymphoid cell lines contain highly active nucleases which degrade mRNA during isolation, the invention uses mRNA preparation methods specifically developed for the isolation of intact mRNA from cells and tissues containing active nucleases. One such method yields total RNA preparations by cell or tissue disruption is an ethanolperchlorate dry be mixture which reduces nuclease action (Lizardi, P. M. gt al., Ang. Biochem., 98: 116 (1979). This method gives intact translatable mRNA.

- [0092] Other methods that have been used for this invention include extraction of cells with lithium chloride plus urea (Auffray, C., and Rougeon, F., Eur. J. <u>Bischem.</u>, 197: 303 (1980)) or guanidine thiocyanate (Chirgwin, J. M. <u>et al.</u> <u>Bischem.</u> 197: by prepare total RNA.
- Biochemistry, 18: 5244 (19/9)) to prepare total rMV.

 [1093] One universal feature of all expressed immunoglobulin light and heavy chain genes and messenger RNAs is the so-called J region (i.e. joining region, see Figure 1). Heavy and light chain J regions have different sequences, but a high degree of sequence homology exists (greater than 80%) within the heavy 3, regions or the 1200 gight chain J regions. The invention provides consensus sequences of light and neavy chain J regions useful in the design of oligonucleotides (designated herein as UIGs) for use as primers or probes for cloning immunoglobulin light or heavy chain makes of the provided of the control of the co
- mouse J₄S sequences (Figure 7).

 [0094] Another utility of a particular UIG probe may be hybridization to light chain or heavy chain mRNAs of a specific constant region, such as UIG-MJK which detects all mouse J₄ containing sequences (Figure 7). UIG design can also include a sequence to introduce a restriction enzyme site into the cDNA copy of an immunoglobulin gene (see Figure 7). The invention may, for example, utilize chemical gene synthesis to generate the UIG probes for the cloning of V regions in immunoglobulin mRNA from hybridoma cells making monoclonal antibodies of desired antigen specificities.

 [0095] A multi-stage procedure is utilized for generating complete V + C region cDNA clones from hybridoma cells light and heavy chain mRNAs. In the first stage, the invention utilizes UIG probes as 'primers' for reverse transcriptase or plementary strand of the primer extended cDNA is then synthesized, and this double-stranded cDNA is cloned in appropriate cDNA cloning vectors such as pBR322 (Subler and Hoffman, Gene, 25: 283 (1989)) or pQ33 (Figure 5). Maniats T, § 8], Molecular Cloning is Abbotatory Manual, Cost Spring Habor Laboratory Publications, New York, page 224 (1982)). Clones are screened for specific hybridization with UIG oligonucleotide probes. Positive heavy and light chain clones identified by this screening procedure are remapped and sequenced to select hose containing N region (1981) clones destricted by this screening procedure are remapped and sequenced to select hose containing N region
- and leader coding sequences.
 [0096] An alternative method is to make cDNA clones using cligo-dT as a primer, followed by selection of light and heavy chain clones by standard hybridization methods.
- 10097] A second stage utilizes cloning of C region gene segments to form heavy and light chain module vectors. In 100 on emethod cDNA clones of human heavy and light chain immunoglobulin mRNA are prepared. These cDNA clones are then converted into C region module vectors by site-frected mutagenesis to place at restriction site at a desired location near a boundary of the constant region. An alternative method utilizes genomic C region clones as the source for C region module vectors.
 - [0089] A third stage of cDNA cloning involves the generation of complete light and heavy chain coding sequence with linked V and C regions. The cloned V region segments generated as above are excised and ligitated to light or heavy chain C region module vectors. For example, one can clone the complete human keaps light chain C region and the complete human gamma, C region, in addition, one can modify a human gamma t region and introduce a termination codon, thereby obtain a gene sequence which encodes the heavy chain portion of an Fab molecule.
- [0099] The coding sequences having operationally linked V and C regions are then transferred into appropriate expression systems for expression in appropriate hosts, prokaryotic or eukaryotic. Operationally linked means in-frame joining of coding sequences to derive a continuously translatable gene sequence without alterations or interruptions of the triplet reading frame.
 - [0100] One particular advantage of using cDNA genetic sequences in the present invention is the fact that they code

continuously for immunoglobulin chains, either heavy or light. By "continuously" is meant that the sequences do not contain introns (i.e. are not genomic sequences, but rather, since derived from mRNA by reverse transcription, are sequences of contiguous exons). This characteristic of the cDNA sequences provided by the invention allows them to be expressible in prokaryotic hosts, such as bacteria, or in lower eukaryotic hosts, such as yeast.

[0101] Another advantage of cDNA cloning methods is the ease and simplicity of obtaining V region gene modules. [0102] The term "non-human" as used in the invention is meant to include any animal other than a human, wherein an immune response can be generated which then leads to usable a cells resulting in corresponding hybridomas or B cell clones obtained by viral transformation and the like. Such animals commonly include rodents such as the mouse or the rat. Because of ease of preparation and great availability, the mouse is at present the preferred, non-human animal. Mouse-mouse hybridomas are thus utilized as the preferred sources for heavy and light chain variable regions. [0103] Preferably, the invention provides entire V and/or C region cDNA sequences. This means that the sequences

code for substantially operable V and/or C regions, without lacking any major structural portions thereof. [0104] The terms "constant" and "variable" are used functionally to denote those regions of the immunoglobulin chain, either heavy or light chain, which code for properties and features possessed by the variable and constant

regions in natural non-chimeric antibodies. As noted, it is not necessary for the complete coding region for variable or constant regions to be present, as long as a functionally operating region is present and available.

[0105] A wide range of source hybridomas are available for the preparation of mRNA. For example, see the catalogue ATCC CELL LINES AND HYBRIDOMAS, December, 1984, American Type Culture Collection, 12301 Parklawn Drive, Rockville, Maryland 20852, U.S.A., pages 5-9 and the ECACC Catalogue, 2nd Edition; PHLS CAMR Porton Down, Salisbury, Wills; SP40JG, U.K. pages 30-35 and 40-46. Hybridomas secreting monoclonal antibodies reactive to a wide variety of antigens are listed therein, are available from the collection, and usable in the invention. Of particular interest are hybridomas secreting antibodies which are reactive with viral antigens, including Dengue complex specific (ATCC HB 114), Dengue type 1 virus (ATCC HB 47), Dengue type 2 virus (ATCC HB 46), Dengue type 3 virus (ATCC HB 49), Dengue type 4 virus (ATCC HB 48), Epstein-Barr receptor (ATCC HB 135), Flavivirus group (ATCC HB 112), hepatitis B surface antigen (ATCC CRL 8017 and 8018), herpes simplex type I (ATCC HB 8068), herpes simplex type II (ATCC HB 8067), influenza virus (ATCC CL 189), influenza A virus, matrix protein (ATCC HB 64), influenza A virus, nucleoprotein (ATCC HB 65), influenza A Bangkok/1/79HA (ATCC HB 66), influenza AWSN NP (ATCC HB 67), SV40 large T antigen (ATCC TIB 115), SV40 large T antigen, C-terminal end (ATCC TIB 117), and SV40 nonvirar T antigen (ATCC TIB 116). Examples of other hybridomas include those secreting antibodies to tumor associated antigens or to human lymphocyte antigens, such as those reactive to human tumor-associated CEA, high mw (ATCC CRL 8019); human tumor-associated alpha-fetoprotein, IgG₁K - (ATCC HB 134); human B lymphocyte HLA-DR, monomorphic,

IgG_{2b} (ATCC HB 104); human T lymphocyte T cell precursors, IgG₁ (ATCC CRL 8022); human T lymphocyte T cell subset, helper, IgG_{2b} (ATCC CRL 8002); T subset, suppressor/cytotoxic, human, IgG₁ (ATCC CRL 8013); T cell subset, suppressor/cytotoxic, human, IgG₂₈ (ATCC CRL 8014); T cells, peripheral, human, IgG₁ (ATCC CRL 8000); T cells, peripheral, human, IgG_{2a} (ATCC CRL 8001); thymocytes. *common, *human, IgG₁ (ATCC CRL 8020).

[0106] These lines and others of similar nature can be utilized to copy the mRNA coding for variable region, using the UIG probes. Of particular interest are antibodies with specificity to human tumor antigens.

[0107] Expression vehicles include plasmids or other vectors. Preferred among these are vehicles carrying a functionally complete human constant heavy or light chain sequence having appropriate restriction sites engineered so that any variable heavy or light chain sequence with the appropriate cohesive ends can be easily inserted thereinto. Human constant heavy or light chain sequence-containing vehicles are thus an important embodiment of the invention. These vehicles can be used as intermediates for the expression of any desired complete heavy or light chain in any appropriate host.

[0108] Among bacterial hosts which may be utilized as transformation hosts, E. coli K12 strain 294 (ATCC 31446) is particularly useful. Other microbial strains which may be used include E. coli X1776 (ATCC 31537). The aforementioned strains, as well as E. coli W3110 (ATCC 27325) and other enterobacteria such as Salmonella typhimurium or Serratia marcescens, and various Pseudomonus species may be used.

[0109] In general, plasmid vectors containing replicon and control sequences which are derived from species compatible with a host cell are used in connection with these hosts. The vector ordinarily carries a replication site, as well as specific genes which are capable of providing phenotypic selection in transformed cells. For example, E. coli is readily transformed using pBR322, a plasmid derived from an E. coli species (Bolivar, et al., Gene, 2: 95 (1977)). pBR322 contains genes for ampicillin and tetracycline resistance, and thus provides easy means for identifying transformed cells. The pBR322 plasmid or other microbial plasmids must also contain, or be modified to contain, promoters which can be used by the microbial organism for expression of its own proteins. Those promoters most commonly used in recombinant DNA construction include the beta-lactamase (penicillinase) and lactose (beta-galactosidase) promoter systems (Chang et al., Nature, 275: 615 (1978); Itakura et al., Science, 198: 1056 (1977)); and tryptophan promoter systems (Goeddel et al., Nucleic Acids Research, 8: 4057 (1980); EPO Publication No. 0036776). While these are the most commonly used, other microbial promoters have been discovered and utilized,

[0110] For example, a genetic construct for any heavy or light chimeric immunoglobulin chain can be placed under the control of the lettward promoter of bacteriophage lambade (P.D.). This promoter is one of the strongest known promoters which can be controlled. Control is exerted by the lambda repressor, and adjacent restriction sites are known. [0111] The expression of the immunoglobulin chain sequence can also be placed under control of other regulatory sequences which may be "homologous" to the organism in its untransformed state. For example, lactose dependent processors IDNA comprises a lactose or lac operon which mediates lactose dispestion by elaborating the enzyme bets-galactostase. The lac control elements may be obtained from bacteriophage lambda pLACS, which is infective for E. [out]. The lac promoter-operator system can be induced by IPTG.

[0112] Other promoter/operator systems or portions thereof can be employed as well. For example, arabinose, coicine E1, galactose, alkaline phosphatase, tryptophan, xylose, tac, and the like can be used. Other bacterial gene expression control elements can be utilized to achieve the expression of immunoglobulin proteins. For example, a gene with a bacterial secretion signal peptide coding region can be expressed in bacteria, resulting in secretion of the immunoclobulin peatide which was originally linked to the siznal peotide.

[0113] Since an immunoglobulin cDNA is comprised only of sequences representing the mature mRNA encoding an antibody protein or its precursor, additional gene expression elements regulating transcription of the gene and processing of the FRNA are required for optimal synthesis of immunoglobulin mRNA. These elements may include spice signals, as well as transcription promoters including includible promoters enhancers, and termination signals. cDNA appression vectors incorporating such elements include those described by Oksayama, H. and Berg, P. Mol. Cell Biol., § 280 (1983); Cepho, C. L. et al., Eugl. 327 : 1053 (1984); and Kauman, R.J. Proc. Natl. Acad. Sci., USA, § 2686 (1985).

[0114] Diffrent approaches can be followed to obtain complete H₂L₂ antibodies.

[0115] First, one can separately express the light and heavy chains followed by <u>in vitro</u> essembly of purified light and heavy chains into complete H₂L₂ IgG antibodies. The assembly pathways used for generation of complete H₂L₂ IgG molecules in cells have been extensively studied (see, for example, Scharff, M., <u>Harvey Lectures</u>. 69: 126 (1974). In <u>vitro</u> reaction parameters for the formation of IgG antibodies from reduced isolated light and heavy chains have been defined by Beychock, S._Gelie of Immunoplobulin Synthesis, Academic Press, New York, page 63, 1979.

[0116] Second, it is possible to co-express light and heavy chains in the same cells to achieve intracellular association and linkage of heavy and light chains into complete H₂L₂ IgG antibodies. The coexpression can occur by using either the same or different plasmids in the same host.

[0117] Co-expression with the aid of secretion signals useful in bacteria, enable fully folded and assembled H₂L₂ immunoolobulins to be obtained.

[0118] Also, preparation of chimeric Fab fragments can be carried out by the methods of the invention.

[0119] The methods described herein can also be used to switch the class of any antibody of a given specificity and class to an antibody of the same specificity but of a different class, whether human or non-human. For example, human IgM antibodies can be transmuted to human IgG antibodies by preparing constructs containing human constant IgG cDNA or genomic sequences, linked to variable human cDNA sequences obtained from a cell producing the original IgM antibody. These constructs are then introduced into appropriate hosts and expressed.

POLYPEPTIDE PRODUCTS

10 [120] The invention provides "chimeric" immunoglobulin chains, either heavy or light. A chimeric chain contains a constant region substantially similar to that present in the heavy chain of a naturel human immunoglobulin, and a variable region having any desired antigenic specificity. The variable region is either from human or non-human origin. [0121] The invention also provides immunoglobulin molecules having heavy and light chains associated so that the overall molecule activities of the provided immunoglobulin molecules and properties. Various types of immunoglobulin molecules are provided rmonovalent, divelent (i.e., with different variable regions), molecules with chimeric heavy chains and non-chimeric light chains, or molecules with variable binding domains attached to peptide moleties carrying desired functions.

[0122] Antibodies having chimeric heavy chains of the same or different variable region binding specificity and non-timeric (i.e., all human or all non-human) light chains, can be prepared by appropriate association of the needed polypetide chains. These chains are individually prepared by the modular assembly methods of the invention. [0123] Chimeric Fab Iragements are also part of this invention.

USES

[0124] The antibodies that can be made by the invention having human constant region can be utilized for passive kinmunization, especially in humans, without negative immune reactions such as erum sickness or anaphylacite kind. The antibodies can, of course, also be utilized in prior art immunodiagnostic assays and kits, in labelled form for in ytto imagine, wherein the label can be a radiocative emitter, or an NMR contrasting agent such as a carbon-13 nucleus,

- or an X-ray contrasting agent, such as a heavy metal nucleus. The antibodies can also be used in vitro localization of anticens by appropriate labelling.
- [0125] The antibodies can be used for therapeutic purposes by themselves in complement mediated lysis or can be coupled to toxins or other therapeutic moletles.
- [0126] Class switching of antibodies is useful when it is desired to change the association, aggregation or other properties of antibodies obtained from cell fusion or hybridoma technology. For example, most human-human monoclonals are of the IgM class, which are known for their ease of reduction and aggregation. Changing such antibodies to other antibody types, such as IgG, IgA, or IgE, is thus of great benefit.
- [0127] Mixed antibody-enzyme molecules can be used for immunodiagnostic methods, such as ELISA. Mixed antito body-peptide effector conjugates can be used for targeted delivery of the effector moiety with a high degree of efficacy and secificity.
 - [0128] Having now generally described the invention, the same will be further understood by reference to certain specific examples which are included herein for purposes of illustration only and are not intended to be limiting unless otherwise specified.

EXPERIMENTAL

15

Materials and Methods

20 Tissue Culture Cell Lines

[0129] The human cell lines GM2146 and GM1500 were obtained from the Human Mutant Cell Repository (Camdon, New Jersey) and cultured in RPM11640 plus 10% feat bovine serum (M. A. Bioproducts). The cell lines \$29.0 and CRL 8017 were obtained from the American Type Culture Collection and grown in Dulbacco's Modified Eagle Medium (DMEM) plus 4.5 g/l glucose (M. A. Bioproducts) plus 10% feat bovine serum (Hyclone, Sterie Systems, Logan, Utah). Medis were supplemented with penicillin/streptomycin (Invine Scientific, Invine, Cellfornia).

Recombinant Plasmid and Bacteriophage DNAs

10 [130] The plasmids pBR322, pl. 1 and pUC12 were purchased from Pharmacia P-L Biochemicals (Milwavkee, Wiscossin). The plasmids pBV2-goe and pSV2-gg were obtained from BRI. (Gailhersburg, Maryland), and are available from the American Type Culture Collection (Rockville, Maryland), pl. 1 as sub-close of the 8.3 Kb Higfill to BamH I fragment of the human IgG1 chromosomal gene. A separate isolation of the human IgG1 chromosomal gene. A separate isolation of the human IgG1 chromosomal gene (Davis, M. et al., Nature, 253. 733) inserted into M13mp10. G-tailed pUC9 was purchased from Pharmacia P-L. DNA manipulations involving purification of polymoration of the Collection and chromosomal gene (Davis, M. et al., Nature, 253. 733) inserted into M13mp10. G-tailed pUC9 was purchased from Pharmacia P-L. DNA manipulations involving purification of polymoration DNA by found tensity centrifugation, restriction anddruclessed digestion, purification of DNA fragments by agarces gel electrophoresis, ligation and transformation of g_ coll were as described by Maniatis, T. et al., Molecular Cholming A. Laboratory Manual, 1 (1982). Restriction endorunclesses and other DNAFNA modifying enzymes were purchased from Boehringer-Mannhelm (Indianapolis, Indiana), BRIL, New England Biolabs (Bewryl, Massaschustetts) and Pharmacia P-L.

Oligonucleotide Preparation

45 [0131] Oligonuclecides were either synthesized by the triester method of loat at [Nucl. Acids Res., 10: 1755 (1982)), or were purchased from ELESEN, Los Angeles, California. Tritylated, deablocked oligonuclecidias were purified on Sephadex-G30, followed by reverse-phase HPLC with a 0-25% gradient of acetonitrile in 10mM triethylamine-acetic acid, pH 7.2, on a C16 uBondapak column (Waters Associated). Detritylation was in 50% acutol acid for 30 min, followed by evaporation thrine. Oligonucleotides were labeled with [gamma-XP]ATP plus 14 polynucleotide kinase.

RNA Preparation and Analysis

[0132] Total cellular RNA was prepared from tissue culture cells by the method of Aufray, C. and Rougeon, F. (Eur. J. Biochem., 107: 303 (1980)) of Chirgwin, J. M. <u>et al. (Biochemistry, 18</u>: 5294 (1979)). Preparation of poly(A)* RNA, methyl-mercury agarose gel electrophoresis, and "Northern' transfer to nitrocellulose were as described by Maniatis, T. <u>et al.</u>, <u>supra.</u> Total cellular RNA or poly(A)* RNA was directly bound to nitrocellulose by first treating the RNA with formaldehyde (White, B. A. and Bancrott, F. C., <u>J. Biol. Chem., 257</u>: 8569 (1982)). Hybridization to filterbound RNA was with nick-translated DNA fragments using conditions described by Margulies, D. H. <u>et al.</u> (Nature, 295: 168 (1982)).

or with ³²P-labelled oligonucleotide using 4xSSC, 10X Denhardt's, 100 ug/ml salmon sperm DNA at 37°C overnight, followed by washing in 4xSSC at 37°C.

cDNA Preparation and Cloning

[0133] Oligo-dT primed cDNA libraries were prepared from poly(A)* RNA from GM1500 and GM2146 cells by the methods of Land, H. <u>et al. [Nucl. Acids Res.</u> <u>9</u>: 2251 (1981)) and Qubler, V. and Hoffman, B. J., <u>Gene</u>, <u>25</u>: 263 (1983), respectively. The cDNA libraries were escreened by <u>in situ</u> hybridization (Manistis, T., <u>supra)</u> with <u>PSP-labelled oligonucleotides using the conditions shown above, or with nick-translated DNA fragments using the conditions of de Lange et al. (Cell, <u>9</u>4: 891 (1983)).</u>

Oligonucleotide Primer Extension and Cloning

[0134] Poly(A): RNA (20 ug) was mixed with 1.2 ug primer in 40 ul of 64mM KC). After denaturation at 90°C for 5 min, and then chilling in ice, 3 units Human Placental Ribonuclease Inhibitor (BRL) was added in 3 ul of 11 mis-HCl, PH 9.3. The oligonucleotide was annealed to the RNA at 42°C for 15 minutes, then 12 ul of .05M DTT, .05M MgCl₂, and 1 mM each of 4ATP, 4dTD 4CTP, and GTP, was added. 2 ul of alpha-²⁴DP (400 Cl/mmol, New England Nuclear) was added, followed by 3 ul of AMV reverse transcriptase (19 units/ul, Life Science).

[0135] After incubation at 42°C for 105 min., 2 ul 0.5 M EDTA and 50 ul 10 mM Tris, 1 mM EDTA, pH 7.6 were added. Unincorporated nucleoides were removed by Sephadex C-50 spun column chromatography, and the FINA-DNA hybrid was extracted with phenol, then with chloroform, and precipitated with ethanic. Second strand synthesis, homopolymer tailing with dGTP or dCTP, and insertion into homopolymer tailed vectors was as described by Gubler and Hoffman, supra.

25 Site-Directed Mutagenesis

35

[0136] Single stranded M13 subcione DNA (1 ug) was combined with 20 ng oligonucleotide primer in 12.5 ul of Hin buffer (7 mM Tris-HC), pH 7.8, 7 mM MgCb, 50 mM NaCl). After heating 10 95°C in a scaled tube, the primer annealed to the template by slowly cooling from 70°C to 37°C for 90 minutes. 2 ul dNTPs (1 mM each), 1 ul ⁹²P-dATP (10 uC)), 1 ul DTT (0.1 M) and 0.4 ul Klenow DNA Poil (20, Boothinger Mannheim) were added and chaine extended at 37°C for 90 minutes. To this was added 1 ul (10 ng) M13 reverse primer (New England Biolabs), and the heating annealing and chain extension steps were repeated. The reaction was stopped with 2 ul of 0.5M EDTA, pH 9, plus 80 ul of 10 mM Tis-HC), pH 7.6, i mM EDTA. The products were phanol extracted and purified by Sephadax G-50 spun column chromatography and ethanol precipitated prior to restriction enzyme digestion and ligation to the appropriate

Transfection of Myeloma Tissue Culture Cells

[0137] A variation of the method of Ochi, A. <u>et al.</u> (Nature, 302; 340 (1983)) was used for proloptest fusion. 50 ml of bactoria at A₅₀₀ of 10.7 were converted to protoplasts by the method of Sandri-Goldin, R. M. <u>et al.</u> (Mol. Cell. Biol. <u>1</u>; 1743 (1981)), then diluted with 20 ml DMEM byte 10% FBS (final volume is 25 ml), Sp20 cells were harvested, pellisted at 2,200 x g, washed, repellisted and resuspended in DMEM at 2-5x109ml. Bacterial protoplasts (10 ml) were mixed with 10x10⁶ 5p20/Coells and pellisted by centrifugation at 4,000 x g at 22°C for 20 min. After pipetting of the supernation, the pellist was suspended in the remaining drop of medium by flicking the tube. 2ml of 10% DMSO, 37% (w/) PEG6000 (Kodak) in DMEM was added dropwise with mixing over 45 sec. Chird 15 sec. 2 ml of 42% PEG6000 in DMEM was added over 45 sec. Complete DMEM (45 ml) was slowly added with mixing. Cells were pellisted at 2500 x g, then washed and celleted thrice.

[0138] The electroporation method of Potter, H. et al. (<u>Proc. Natl. Acad. Sci., USA</u>, <u>Bi: 7161</u> (1984)) was used. After transfection, cells were allowed to recover in complete DMEM for 48-72 hours, then were seeded at 10,000 to Sci. Cells per well in 95-well culture plates in the presence of selective medium. G418 (GIBCC) selection was at 0.8 mg/ml, mycophenolic acid (Calbicchem) was at 6 ug/ml plus 0.25 mg/ml xanthine, and HAT (Sigma) was at the standard concentration.

Assays for Immunoglobulin Synthesis and Secretion

[0139] Secreted immunoglobulin was measured directly from tissue culture cell supernatants. Cytoplasmic protein extract was prepared by vortexing 1x10⁹ cells in 160 u of 1% NPA0, 0.15 M NaCl, 10 mM Tris, 1 mM EDTA, pH 7.6 at 0°C, 15 minutes, followed by centrifugation at 10,000 x g to emoreus insolutele debris.

[0140] Double antibody sandwist [182] (ISA (Volter, A et al., in <u>Manual of Clinical Immunology</u>, 2nd Ed., 2nd Ed., Rose, N. and Friedman, H., pp. 389-371, 1890) using affiting purified antibarea was used to dest specific immunology. 2nd Ed., Service of attention of human logs, the plate-bound antiserum is goat anti-human IgG (KPL, Gaithersburg, Maryland) at 11/100 (dilution, while the proxidase-bound antiserum is goat anti-human IgG (KPL Tago, Buringame) at 11/4000 dilution. For detection of human immunoglobulins is goat anti-human IgG (KPL Tago, Buringame) at 11/4000 dilution. For detection of human immunoglobulins is goat anti-human IgG (KPL and Immunoglobulins is goat anti-human IgG (KPL a

[0141] Antibodies binding hepatitis B surface antigen were detected using a commercial (Abbott, AUSAB) assay.

EXAMPLES

τO

15

35

[0142] The following examples, of which Example VII examplifies the invention, show cDNA cloning for the modular assembly of antibody genes as used for the preparation of chimeric antibodies each having a human constant region and a non-human variable region. These examples outline the step-by-step process of preparing the chimeric antibod-

EXAMPLE-I; Human Antibody Constant Region Gene Modules and cDNA Expression Vectors

(1) Preparation of cDNA Clones, and Vehicles Containing Same, for Heavy Chain Human Constant Region

6 [0143] The cell line GM2146 was used as the source in mRNA preparation and cDNA cloning. This cell line secretes IgG, (Simmons, J. G. <u>et al.</u>, <u>Scand. J. Immunol.</u>, <u>14</u>: 1-13, 1981). Tests of this cell line indicated that it secretes IgA as well as IgG.

[0144] The cell line was cloned, and results indicated that five of aix subclones secreted [aG only, while one of aix subclones secreted [aG only, Pbc/lA**] FNA was prepared from the poly(A)* FNA was prepared from the poly(A)* FNA by the method of Gubler, U and Hoffman, B. J. Gane, 25 253-258 (1983), An initial plating of the cDNA transformed into E. Coll strains HB101 and RFIV yielded a total of 1500 colonies, which were screened by hybridization to a <u>Highdill to Bamill tragment</u> of a genomic clone of human [gG1 (pHu-gamma-1). Four positive clones were found. A fragment containing the CFI acoding region of one of these clones, pGMH-16 (Figure 4), was used to rescreen the original library plus a new transformation of approximately 5000 colonies. Two of the largest clones, pGMH-16 and pGMH-15, were analyzed by restriction enzyme digestion (Figure 4). Both clones contained the entire constant region of human [gG1, tilhough it was discovered that pGMH-16 had deleted approximately 1500 base pairs of pBR322 DNA apparently without affecting the IgG1 DNA sequences.

[0145] Clone pGMH-6 provided the IgG1 constant region module in the construction of cloning vectors for heavy chain variable region cloning.

(2) Preparation of cDNA Clones, and Vehicles Containing Same, for Light Chain Human Constant Region

[0146] A human cell line (GM1500) producing IgG2K was selected for the initial cloning phase. Poly(A)* RNA prepared from GM1500 is active in <u>in yitle</u> translation using rabbit reliculocyte extracts. A cDNA library was prepared from this RNA by the method of Land <u>et al.</u>, <u>Nucl. Acids Ras.</u> 9: 2251-2266 (1981), utilizing <u>Konl</u> digested and d6-failed pC23 as the cloning vector (Figure 5). This vector contains <u>Bg1</u>II, <u>Kpn1</u> and <u>Ss1</u> sites inserted between the <u>Bam</u>HI and <u>Ss1</u> sites in <u>Ss1</u>

[0147] In order to Identify the cDNA clones generated from GM1500 RNA which correspond to light chain mRNA, a DNA probe, UGH-HIK, was synthesized and purified. The UGH-HIK oligonucleotide has the sequence 5'AGCCAC AGT-TCGTTT-3', and is designed to hybridize to all functional human keppa mRNA species at the J-C junction. This probe was used to prime cDNA synthesis on GM1500 RNA in the presence of olidoxynucleotides and reverse transcriptase. From 1.2 up of total GM1500 poly(A)+ RNA was used in this experiment, the entire J sequence and some of the region was read, demonstrating that (1) GM1500 RNA is intact, (2) the kappa probe is of the correct sequence, and (3) GM1500 light chain mRNA contains J₄4 sequences.

[0148] cDNA clones positive for hybridization to the light chain probe were selected. Since the probe hybridizes to the J-C junction, the most important point was to determine if the clones had complete constant region sequence in addition to the J region.

[0149] Insert sizes for the two largest <u>kappa</u> cDNA chones were 0.6 and 0.9 kb, restriction enzyme mapping indicated that the entire constant region coding sequence was present in both clones (Figure 6). The human <u>kappa</u> cDNA clone pk2.3 was used to make the light chain constant region vector plNa2001 by inserting the <u>Sau</u>9A fragment comprising the <u>human kappa</u> constant and J regions into the Bell site of pBR325 (Figure 6B).

[0150] A variant of the human <u>kappa</u> cDNA clone was made by placing a <u>Hind</u>III site in the J region. This was carried out by in vitro mutagenesis using a J_KHINDIII oligonucleotide primer (Figure 7c). The resultant plasmid is pGML60.

[0151] A vector, pINS2003, was constructed for the transfer and expression of cDNA sequences in mammalian calls (Figure 10). This vector was constructed from pUC12 and two plasmids containing SV40 sequences, pL1 provides an SV40 early region promotier and an SV40 late region splice sequence, pSV2-neo sequences provide a selectable marker for mammalian cell transformation and SV40 polyadenylation signal sequences. pUC12 provides a multiple coning site for cDNA insertion.

[0152] The pING2003 vector has several useful restriction sites for modifications. These include a <u>HindIII</u> site useful for the insertion of afternate promoter sequences. This vector is useful in the expression of cDNA genes in mammalian cells.

10 Addition of Enhancer Element to pING2003

25

[0153] Immunoglobulin enhancer elements have been shown to enhance transcription of genes in their vicinity in stably transformed mouse myeloma cells by several hundred fold (Gillies, S. D. et al. Cell, 33: 729, 1983). To facilitate expression of the mouse-human immunoglobulin genes in mouse myeloma cells, the mouse immunoglobulin heavy chain enhancer element was added to the cDNA expression voter piN26203 (Figure 10). The mouse heavy chain enhancer region DNA was isociated from an M13 subclone of mouse heavy chain enhancer region DNA was isociated from an M13 subclone of mouse heavy chain enhancer region DNA was isociated from a Sall plus Eccell digestion of this subclone was modified with pindIII likers and inserted into the InfindII side of piN262003, restulting in the new cDNA expression vector piN26203E. This vector is useful in the efficient expression of cDNA genes in mammalian cells, particularly mouse myeloma or hybriddna cell lines.

EXAMPLE II: Human-Mouse Chimeric Anti-HBsAG Antibody Chain

(1) Preparation of cDNA Clones and Vehicles Containing Same, for Heavy Chain Mouse Anti-HBsAg Variable Region.

[0154] The cell line CPL8017 was obtained from the ATCC and subcloned. Subclones were grown and tested for mouse IgG anti-hepatitis B binding activity using a commercially available anti-HBsAg detection kit. Three positive subclones were found; Poly(A)* RNA was prepared from one of these subclones, and was fractionated on a methyl-mercury agences gel. The RNA contained intact light chain and heavy chain mRNA's as interred from specific hybridization to kagepa Uid-MNA primer, and to the mouse heavy chain UIG-MNB prove (see Figure 7). In addition, the UIG-MNB primer was used for specific priming of anti-HBsAg poly(A)* RNA in a dideoxy sequencing reaction. Sufficient sequence was read to show that a major kagep RNA of the anti-HBsAg cell fine contains the Ju2 sequence.

[0158] The conditions for variable region cDNA synthesis were optimized by using heavy and light chain UIB primers on anti-HBsAg poly(A)* RNA. Dideoxy chain stension syspriments demonstrated that the mouse UIG-AUJK primer and UIG-JH3 primer correctly primed kappa and heavy chain RNAs. When the reverse transcription was carried out in the absence of dideoxynucleotides, the main product using he lagogs uIG-AUJK primer was a 410±20 nucleotide fragment, while the main product using the heavy chain IUG-JH3 primer was a 430±30 nucleotide fragment. These correspond to the expected lengths of the variable and 5' untranslated regions of kappa and heavy chain immunopiculum RNAs. The conditions for the optimal priming of poly(A)* RNA from CRL8017 calls should work wall for poly(A)* RNA isolated from any cell line producing a monoclonal antible of the contractions.

[0156] After determining optimal conditions for priming hybridona mRNA with oligonucleotide primers, two oligonucleotides were designed and used for heavy chain V region DNA synthesis. These two oligonucleotides are IUGA.MJHBSTEII(13) and IUGA-MJHBSTEII(13) primer was 13 bases to make the analogous position at the latter's J region. In this case, the primer had a single G io U mismatch with the mouse mRNA sequence that uses the J₁/3 coding sequence. The IUGA-MJHBSTEII(13) primer was 13 bases in a make the make the make the MJHBSTEII(13) primer was 13 bases in make the make the MJHBSTEII(13) primer was 13 bases in make the MJHBSTEII(13) primer was 13 bases in make the make the MJHBSTEII(13) primer was 13 bases in make the MJHBSTEII(13) primer was 14 bases in make the MJHBSTEII(13) primer was 14 bases in make the MJHBSTEII(13) primer was

[0157] These two primers and the J_h3 coding sequences are shown in Figure 8. The first strand cDNA products made via the 13-mer BatEll and the 21-mer J_h3 primers included bands of approximately 430 nucleotides, which represented the entire V_h1 region. Under the standard priming conditions used, the priming efficiency of the 13-mer BstEll was much less than that of the 21-mer J_h2. Accordingly, a CDNA library was generated from the first strand synthesis from each of these primers, using the method of Gubber and Hoffman, supra.

[0158] First, the 21-mer J₂5 library was screened with the 21-mer J₂5 oligonucleotide. Filter hybridization was done at 30°, overnight, according to de Lange, T₂ gt_1, Cell_32. Sel pol 90 (1993). The filters were then washed at 51° in 6 x SSC, 0.1% SDS, Five colonies were selected. The largest had an insert of approximately 450 bp. More significant it contained three restriction sites predicted from the known J₂3 sequence, which are present upstream of the primer.

sequence. This clone, p.13-11, was sequenced using the J₂3 primer by the chain-termination method (Wallace, R. B. et al., <u>Gene.</u> 16: 21-26 (1981)). The sequence obtained has the remaining J₄5 coding segment. Just upstream a 13-nucleotide segment matched to a published D segment sequence (Dsp 2-2) (Kurosawa, V. <u>et al.</u> J. <u>Exo. Med.</u> 155; 201 (1982), and Tonegawa, S. <u>Nature.</u> <u>302</u>: 575 (1983)). A norapeptide pradicted from this area showed characteristic homology to the published mouse heavy chain to subgroups at amno sed residues 8 to 8 4, comparish the F3 of heavy chain molecules. Plasmid pJ3-11 represented a rearranged VDJ sequence, and apparently contained the anti-hepatilit V₁ sequence produced by the cell line.

[0.59] In order to isolate a V_{II} region cDNA clone that had a <u>Bat</u>EII site in the J region, an <u>Alul</u> to <u>Sau961</u>, 255 nucleotide long, probe from pJ3-11 was next used to screen the cDNA library generated from the 13-mer <u>Bat</u>EII primer. Six positive clones were isolated. The largest, pBs13-1, was further analyzed. The insert was 280 nucleotides long and its restriction map agreed with that of pJ3-11 except for the introduced <u>Bat</u>EII site. Figure 3 illustrates how these two inserts were recombined to generate pMY-EG-13, a. V_{II} clone with the module-pioling <u>Bat</u>EII site. Three additional V_{II} cDNA clones were isolated from a cDNA library generated from the 21-mor oligonucleotide UIG-MUH3BSTEII primer containing a <u>Bat</u>EII site. Three additional

(2) Preparation of cDNA Clones, and Vehicles Containing Same, for Light Chain Mouse Anti-HBsAg Variable Region

15

[0160] Since the Ju2 sequence is present in mRNA propered from the anti-hepatitis hybridoms cell line, the oligonuclocide utils (JSCBGUI (Figure 7B), was designed to introduce a <u>Boll</u> list into the Ju2 region. Digestion with Bull would then allow direct insertion of a V_x cDNA coding region into the <u>Boll</u> site of the previously noted human C_x vector, piNC2001. This insertion would result in the procise joining of a mouse variable region segment (including the 1 region) to a human <u>sepage</u> constant region segment, each in the proper coding frame and with no alteration in amino acid sequences for either mouse variable or human constant region.

[0161] The JKZBGLI oligonucleotide was used to prime anti-HBsAg mRNA to form a cDNA library as for heavy chain, <u>supra</u>, in pUC9. The cDNA was size-selected by polyacrylamide gel electrophoresis prior to cloning, and 80% of the cDNA clones were shown to have insert sizes between 300 and 750 nucleotides in length. Replica filters of this library were screened with two oligonucleotides, the original primer and a second probe complementary to J₄2 sequence 5 to the original primer.

[0162] It was discovered that the anti-hepatitis B monoclonal cell line CRL 8017 secrates immunoglobulins with at least two different light chains, One of them is derived from the myeloma NS-1, which was used as a fusion partner in generating the anti-hepatitis B cell line. Since NS-1 is derived from the myeloma MOPC21, the possibility was investigated that MOPC21 \(\text{V}_m RNA may be present in the V_C cONA library from the anti-hepatitis monoclonal cell line. Indeed, one cDNA clone (p6D4B) analyzed has an identical restriction enzyme map to that of MOPC21 \(V_C CDNA, accept for the inserted Bulls lise. \)

© [0143] Two conclusions can be drawn from these results. The first is that it is possible to effectively use an oligonuclecidie to includuce a restriction enzyme site while cloning a V_L region from a hybridome coll line. The second is that one must carefully monitor hybridoma cell lines for the presence of multiple V region sequences, only one of which is the delered sequence.

[0164] In order to further characterize the <u>kapps</u> light chain J regions present in the cell line mRNA, poly(A)+ RNA was bound to nitrocellulose by the formatidehyde "Dot blot" procedure of White and Bannorth, <u>J. Biol. Chem., 527</u>: 9599 (1982). The RNA was hybridized to ³²⁹-labeled oligonucleotide probes specific for each functional <u>kapps</u> J region. These probes are shown in Figure 79 as the Ulig probes SIA*, MIX, SIA*4, and SIA*5. The results haved that the mRNA hybridized strongly to both MJK and SIA*6 oligonucleotide probes, inclicating that both Juk² and Juk² sequences we present. Since Jug² mRNA had been previously identified as the one derived from the parental hybridoma partner NS-1, it was concluded that the Juk² mRNA encoded the anti-hepatits binding specificity of the CRIL 8017 cells:

NS-1, it was concluded that the J₄4 minNA encoded the anti-nepatities binding specificity of the C+L-1 of Testis. [D165] Two different CDNA libraries were ceremend to isolate V region clones encoding J₄4 sequences. The first was primed by JK2BCLII, gupra. The second was made by using the oligonucleotide primer, JK4BCLII, which is specific for J₄4 miNNA and introduces a Bigli stall into the J region of cloned V regions. The JK4BCLII primer was used to prime first strand cDNA synthesis to construct a cDNA library by the same method used to construct a JK2BGLII primed cDNA library except that CDNA was not size selected prior to cloning.

[0166] Figure 7B tabulates the mismatches that each primer has with other functional mouse <u>kappa</u> J region sequences. Note that U_AL has five mismatches in 21 nucleotides when compared with the JK2BGLII primer, and 3 in 23 with the JK4BGLII primer.

[0167] Both libraries were screened for V region clones containing J_K4 sequences by hybridizing to an oligonucleotide probe specific for J_K4 sequences (5JK4). The results of this screen are shown in Table 1.

Table 1*

Library	Probe Specificity		
	J _K 2	J _K 4	
JK2BGLII	2% (30/1500)	0.15% (2/1500)	
JK4BGLII	N/D	3.5% (31/875)	
* Descente se of closes containis	n 1 2 or 1 4 coguence plus a V region	The probes used were the olinopucle.	

*Percentage of clones containing 3_x2 or 3_x4 sequence plus a V region. The process used were the diigonucleotide 5JK4 (J_K4 specificity, Figure 7) and p6D4B, which contains the NS-1 (MOPC21) V region sequence. N/D, not done.

[0168] Saveral J₄ AV region cDNA clones isolated from both libraries were characterized. These clones have identical restriction anzyme maps, including the engineered <u>Bell</u> list in evaluting from the cligonuclectide primed cDNA cloning procedure. The restriction map and sequence of one clone, pV17, show that pV17 contains V region gene sequences. [0169] These results show that that JA/2BGLI primer could correctly, although in efficiently, prime <u>J</u>4 mRNA sequences. Since the JK2BGLI primer had less mismatches with any other <u>J</u>4 region mRNA than with <u>J</u>4 mRNA (Figure 78). It is expected that the other JK mRNAs can be primed at the correct location with better efficiency using the JK2BGLI primer. Thus, efficient cDNA cloning of any functional mouse <u>kapps</u> V region may be obtained by using a mbture of the JK2BGLI and JK4BGLI Unimers.

[0170] The placement of a <u>Bull</u>l site into the J region during cDNA cloning of the V regions allows joining of the cloned mouse V region gene module to the human <u>kapon</u> constant region gene module (Figure 9B).

[0171] After the aforementioned experiments were carried out it was found that the cDNA clone pV1 lacked a complete 5" coding region. Nucleotide sequencing showed that the A of the initiation codon ATG was not copied in pV17. This was not a random cDNA cloning artifact because two other cDNA clones had the same defect. Two approaches were devised to obtain a light chain gane with a complete 5" coding region.

(IT2) First, a new CDNA library was constructed by first printing with an oligonucleotide (5-ATATTGCT-GATGCT CT-3) complementary to pV17 sequences 155 bases from the 5 and From this library, clones hybridizing to a pV17 DNA tragment probe were selected, and some of these new cDNA clones have the initiator ATG plus about 20 nucleotides of 5' untranslated region. One of these clones, p2-12, supplies a 5' untranslated region of 23 nucleotides and a compliet ATG initiator codon. When p2-12 was combined with pV17 derived sequences, a variable region with a

complete 5' and was formed (pING2013E).

[0173] Second, elled-frected mutagenesis on the existing light chain clone was used to simultaneously remove the poty-G tract and place a ribosome recognition sequence adjacent to the initiator ATG. The Pati fragment from pVT was subcined into M13mpl E.A on digenulcediate (V11-VM). 5'-G1TGCAGCTAGAGTAGAGTTCC AGGTTC-S vas then used as a primer to mutate the pVT7 sequence to include a Sall site and an initiator ATG into the pVT7 sequence.

The resultant plasmid pVT7-WM provided an atternate mouse variable region for joining to human constant region.

modules.

[0174] The complete nucleotide sequence of the variable region from pV17 was then determined. The sequence shows that pV17 contains a V_{K-V} junction region, containing several conserved amine acids, and the hybrid A₂Z₁J₄ region formed by priming the J₄Z RNA with the UIG-JK2BGLII oligonucleotide. However, the V_K region in pV17 is non-functional, because the V_K and J_K regions are not in the same coding frame. Translation of the pV17 V region would have result in an abnormal immunoglobulin light chain where the J region is translated in an incorrect frame. This defect may be caused by aberrant V-J joining, resulting in a non-functional <u>kappa</u> mRNA, as has been observed by Kelley, DE, et al., Mol. Cell Biol. 5-160-0175 (1980).

[0175] Since the pV17 V region encodes an abnormal immunoglobulin, it is highly unlikely that this light chain is part of a functional anti-hepatitis entibody molecule. These results show the importance of monitoring hybridoria cells for the presence of multiple RNA species encoding V regions, only one of which is the desired sequence.

[0176]. Further screening of CRL 8017 CDNA libraries was done to search for V_x CDNA clones which are not more either of the two V_x CDNA classes found so far (McPC21-pD6A), pv17]. First an oligo-dT primed cDNA library made from CRL8017 RNA was screened with a DNA fragment probe specific for the <u>kapps</u> constant region, and separately with probes specific for MCPC21 and pv17 V_x regions. A DNA clone (giElS-41) that contains the <u>kapps</u> constant region, but has a different V_x region than that of McOPC21 or pV17 was discovered. This method of screening digo-dT primed cDNA libraries is a useful alternative to oligonucleotide screening of CDNA libraries lose abscause nick-transled probes of high specific activity are used. Also, this method allows the simultaneous isolation of several classes of V region clones, such as all V_x clones, by appropriate probe choice. Second, the UGI-X/B2DLI-primed cDNA library made from CRL 8017 RNA was screened with the UGI-SJRZ digonucleotide probe (see Figure 7). A new class of V_x cDNA clones was found whose members are homologous to pt618-014 and hydridze to the UGI-SJRZ probe, but

to a MOPC21 V_K probe. The restriction endoruclease site maps and nucleotide sequences of these clones also differ rom MOPC21-homologous V_K cDNA clones from CHB.017 cBis. These clones, however, have an aberrant Vollet which results in a nonfunctional mRNA, and appear to be identical to one described by Cabilly and Riggs (Gane, 40: 157.11985).

[0177] It was therefore concluded that the anti-hepatitis B cell line CRL8017 has at least three classes of V_K mRNA corresponding to the above described cDNA clones p6D4B (MOPC21), piE9L, and pV17. The piE9L and pV17 clones are derived from mRNA from aberrantly rearranged Kappa genes, while the p6D4B clone is derived from the parent hybridoma fusion partner NS-1. None of these clones appear to encode the desired anti-hepatitis light chain.

(3) Preparation and Expression of Heavy Chain Containing Human Constant/Mouse Variable Regions

[0178] The V region sequences in pMVHCa-13 were joined to the human IgG1 constant (C) region clone pGMH-6. Due to the presence of a second BatEll site within the IgG1 CH1 region of pGMH-6, a multi-step ligation was required. First, the 220 nucleotide BatEll it regment from the J-CH1 region of pGMH-6 was ligated to the 1100 nucleotide IgG region BatEll to BatHl I fragment of pGMH-6. In a separate ligation, the 420 nucleotide BatEll to BatHl I fragment of pMVHCa-13, which comprises the mouse V region, was pioned to a cell intestine phosphatase treated BatHl plant of working the comprises the mouse V region, was pioned to a cell intestine phosphatase treated BatHl plant over the combined, ligase was added, and the products were transformed into HB101, resulting in the chimeric mouse V-human C close pMVHCa-14, (Figure 9A).

[0179] The V region of the hybridheavy chain gene in pMVHCc-24 was further analyzed by partial sequence analysis. This analysis showed that the clened V region contained a D sequence within matches a known D sequence, DSP2.2 (Kurosawa and Tonegawa, <u>supra</u>). The sequence also predicted a 19 amino acid leader peptide similar to known mouse V heavy chain leader peotide sequences and a 5 untranstated region of at least 3 nucleotides.

[0180] The BamHI fragment containing the mouse-human hybrid heavy chain gene of pMVHCc-24 was cloned into BamHI digosted pIN82003E vector, resulting in the expression plasmid plN8206E (Figure 11). The pN82008E indigoration of the mouse-human chimeric immunoglobulin gene in 8 mythodic does because of the presence of the mouse heavy chain enhancer region.

[0181] A modification of the chimeric heavy chain gene present in pMVHCc-24 was done to provide an alternate heavy chain gene which lacks the oligo-dc region preceding the initiator ATG. The pING2012E and pING2008E vectors are identical except for the nucleotides immediately preceding the ATG, as shown in Figure 12.

Vei [182] Bacteria harboring the pING2005E and p5V2-neo plasmids were converted into protoplasts by the method of Sandri-Codin, R. M. et al., Mol. Cell. Biol. j. 743 (1981). The protoplasts were then separately fused to SP20-ApJ4 hybridoma cells (ATC CFIL. 1581) by freatment with polyethyleneplycol (Chrit. A. <u>et.al.</u>, Nalurg. 302: 340, 1983). The used colls were allowed to recover for 72 hours in complete nedium before plating at 10 0,000 or 50,000 cells per well in a 96-well lissue culture plate. The cells were selected with G418 at 0.8 mg/ml for two weeks, when growth in some wells was clearly evident. Under these selection conditions, Sp20 cells were completely killed within 47-days by G418. Only cells which have integrated and expressed the neo gene present in the vectors will grow under G418 selection. The number of wells positive for growth by these integrative transfectants are shown in Table 2.

Table 2*

40

Strain/Plasmid	10,000 cells/well	50,000 cells/well
MC1061/pING2006E	3 (13%)	12 (50%)
MC1061/pSV2-neo	7 (29%)	4 (17%)
MC1061/none	0	0
* Percentage of w	ells showing positiv	e growth out of 24

[0183] Cells transfected with pINS2006E and pSV2-neo were tested for immunoglobulin gene expression at the RNA and protein level. Total cell RNA was prepared from transfected cells, bound to nitrocellulose and hybridized to RNA was prepared from transfected cells, bound to nitrocellulose and hybridized to RNA was prepared to represent the properties of the mouse-human hybrid heavy chain gene. Two clones were found which have a strong signal, representing expression of the gene at the RNA level. The amount of total cellular RNA hybridizing to the mouse-human probe appeared to be approximately 1/10 the level of heavy chain RNA in the original hybridoma cells. This probably represented about 176 of the total mRNA of the transfected cell.

[0184] The transfected mouse cells were also tested for production of cytoplasmic human heavy chain protein by an ELISA assay, It was found that 3 out of 7 pilos/2006 transfected cell lines produced detectable levils of human heavy chain protein. The mouse cell transformant producing the most mouse-human heavy chain protein pave a signal in the ELISA assay comparable to that of a 17/100 diblint of a human B cell little producing intext human immunoglobulin

IgG1. This modest level of detected mouse-human heavy chain protein may be due to several factors, including instability of heavy chains in the absence of light chains in hybridoma cells, or incorrect processing of the chimeric gene transcript.

5 (4) Gene Amplification of the Integrated Chimeric Gene

[0165] Southern blot analysis showed that multiple copies of the pNIQ2006E DNA sequences were integrated in tandern in the mouse genome. Restriction enzymes <u>Agal and Bull</u> blot eleave point/2006E eingly, in the transformant, 2AE9, a band, from an <u>Agal</u> or <u>Bull</u> digestion, of the expected size (8.2/bt) was found to hybridize to the human C gamma is sequences (data not shown). An <u>a Bam</u>th band of the correct size (1.6/bt) was found to hybridize to the human as well as the IE9 V_H sequences. Gene-copy iltration experiment (Fig. 14) indicated that there are about 5 copies of pING2006E in the 2AE9 genome. That fact that only a single band was detected in the <u>Agal or Bull</u> indicates that these individual copies are in a tendemly arranged array. A set of double digestions showed that nNG2006E sequences suffered no rearrangement in their introduction into the mouse DNA (data not shown).

[0188] We next transfected the 2AE9 cells with a plasmid that contains a different selectable marker, the and selected clones growing out in DMEM-HAT. One clone, 2BHIO, has about 38 ng soluble human gamma I protein per 10° cells. Southern analysis showed that 2BH10 has about 30 copies of pING2008E (Fig. 14). They were amplified from the 5 copies in 2AE9 without rearrangement of the DNA sequences. (Compare the 2AE9 panel to the 2BH10). SI data (data not shown) revealed that this increase in template led to a higher amount of 19G gene transcripts. We

believe that these sequences were co-amplified with contiguous cellular sequences as a result of the second selection.

EXAMPLE III: A Human-Mouse Chimeric Antibody with Cancer Antigen Specificty

(1) Antibody L6

[9187] L6 monoclonal antibody (MAb) was obtained from a mouse which had been immunized with cells from a human lung accinome, after withis aplean cells were hybridized with NS-1 mouse myelome cells. The antibody brids to a previously not identified carbohydrate antigen which is expressed in large amounts at the surface of cells from most human carcinomes, including lung carcinomes (adeno, squarmous), breast carcinomes, collo carcinomes and oversine carcinomes, while the entigen is only present at Irace levels in normal cells from the adult host. MAb L6 is an IgG2a and can medical entitle or the presence of human peripheral blood eluckocytes as a source of effector cells, or as to type L6 positive tumor cells, and it can yes L6 positive tumor cells in the presence of human serum as a source of economic control of the presence of human serum as a source of complement; the lysis is detected as the release of ⁵¹Cr from isbelled cells over a 4 hour incubation period. MAb L6 can localize to L6 positive tumors exorteransplanted onto nude mice, and it can inhibit the outgrowth of such tumors. MAb L6 is described in <u>Cancer Res. 46</u>:3917-3923, 1986 (on MAb specificity) and in Proc. NALL Acad. Sci. 82:3759-7033, 1986 (on MAb function).

(2) Identification of J Sequences in the Immunoglobulin mRNA of L6.

- [0188] Frozen cells were thawed on ice for 10 minutes and then at room temperature. The suspension was diluted with 15 ml PSB and the cells were centifuged down. They were resuspended, after washes in PSB, in 16 ml 3M LICI, 6M urea and disrupted in a polytron shear. The preparation of mRNA and the selection of the poly(A*) fraction were carried out according to Auftra, C. and Rougeon, F. Eur. J. Bischem. 107:303, 1980.
- [0189] The poly (A*) FINA from L6 was hybridized individually with labeled J_H1, J_H2, J_H3 and J_H4 oligonucleotides ander conditions described by Nobrega et al. Anal. Biochem 131:141, 1983). The products were then subjected to electrophoresis in a 1.7% agarose-TBE gel. The gel was fixed in 10% TCA, bottled dry and exposed for autoradiography. The result showed that the L6 v_H contains J_H2 sequences.

[0190] For the analysis of the \(\frac{\chi_A}{\chi}\) mRNA, the dot-holt method of White and Bancroft J. Biol. Chem. <u>257</u>:859, (1982) was used. Poly (A*) RNA was immobilized on nitrocellulose filters and was hybridized to labeled probe-oligonucleotides at 40° in 4xSSC. These experiments show that L6 contains \(\frac{\chi}{\chi}\) is equences. A faint hybridization to \(\frac{\chi}{\chi}\) was observed.

(3) V Region cDNA Clones.

[0191] A library primed by oligo (dT) on LE poly (A*) RNA was screened for kappa clones with a mouse C_v region probe. From the L6 library, several clones were isolated. A second screen with a 5' J_oS specific probe identified the L6 (J_oS) light-chain clones. Heavy chain clones of L6 were isolated by screening with the J_oZ oligonucleotide. [0192] The heavy and light chain genes or gene fragments from the ONA clones, pH3 6a and pL3-12a were inserted of the M3 bacteriophage vectors for nucleotide sequences and the variable.

region of these clones were determined (FIGURES 15 and 16) by the dideoxy chain termination method. These sequences predict V region amino acid compositions that agree well with the observed compositions, and predict peptide sequences which have been verified by direct amino acid sequencing of portions of the V regions.

[0193] The nucleotide sequences of the cDNA clones show that they are immunoglobulin V region clones as they contain amino acid residues diagnostic of V domains (Kabat et al., Sequences of Proteins of Immunological Interest; U.S. Dept of HHS, 1983).

[0194] The L6 V_H belongs to subgroup II. The cDNA predicts an N-terminal sequence of 24 amino acid residuas identical to that of a known V_H (45-165 CRI; Margolies <u>et al. Mol. Immunol.</u> 18:1065, 1981). The L6 V_H has the J_H2 sequence. The L6 V_H is from the V_K-(Kprl family (Nish <u>et al. Proc. Nat. Acc. Sci. USA 82</u>:399, 1985), and uses J_H5. The cloned L6 V_T predicts an amino acid sequence which was confirmed by amino acid sequencing of peptides from the L6 light chain corresponding to residues 18-40 and 80-96.

(4) In Vitro Mutagenesis to Engineer Restriction Enzyme Sites in the J Region for Joining to a Human C-Module, and to Remove Oligo (dC) Sequences 5' to the V Modules.

[0195] Both clones generated from priming with oligo (dT) L6 V_A and L6 V_A need to be modified. For the L6 V_K , the J-region mutagenesis primer J_K^{+} Hmill, as shown in FIGURE 178, was utilized. A human C_K module derived from a CDNA clone was mutagenized to contain the $\frac{1}{M}$ Hmill sequence (see Figure 17A). The mutagenesis reaction was performed on M13 subclones of these genes. The frequency of mutant clones ranged from 0.5 to 1% of the plaques obtained

obtained.

[0186] It had been previously observed that the oligo (dC) sequence upstream of the AUG codon in a V_H chimericgene interferes with proper splicing in one particular gene construct. It was estimated that perhaps as much as 70% of the RNA transcripts had undergone the mis-splicing, wherein a cryptic 3' splice acceptor in the loader sequence was used. Therefore the oligo (dC) sequence upstream of the initiator. AUG was removed in all of the clones.

[0197] In one approach, an oligonucleotide was used which contains a <u>Sell</u> restriction elte to mutagenize the L6 V_K clone. The primer used for this oligonucleotide-directed mutagenesis is a 22-mer which introduces a <u>Sell</u> site between the oligo (cd) and the initiator <u>met</u> coden (FIGURE 19).

[0198] In a different approach, the nuclease BAL-31 was used to chew away the oligo (dC) in the L6 V₂ clone pH3-6a. The size of the deletion in two of the mutants obtained was determined by nucleotide sequencing and is shown in PicuIRE 17. In both of these mutuants (delta 4 and delta 21), all of the oligo (dC) 5' to the coding region were deleted. [1919] These clones were then modified by oligonucleotide-directed mutagenesis with the MUH2-Agal primer (FIG-URE 17). This 31-base primer introduces an Agal site in the mouse C₄ gene at a position analogous to an existing Agal site in human Cgamma 1 cDNA gene module. The primer introduces the appropriate codons for the human C gamma 1 gene. The chimeric heavy chain gene made by joining the mutagenized mouse V₄ gene module to a human 5 C₄ module thus encodes a chimeric pricate which contains no human amino acids for the entire V₄ region.

[0200] The human C gamma 1 gene module is a cDNA derived from GM2146 cells (Human Genetic Mutant Cell Repository, Newark, New Jorsey). This C gamma 1 gene module was previously combined with a mouse V_H gene module to from the chimeric expression plasmid pING2012E.

40 (5) L6 Chimeric Expression Plasmids.

15

[0201] L6 chimeric heavy chain expression plasmids were derived from the replacement of the V_s module plnCs2012E with the V_s modules of mutants delta? 1 and delta 4 to give the expression plasmids plnCs2111 and plnCs2112 (FIGURE 17). These plasmids direct the synthesis of chimeric L6 heavy chain when transfected into mammalian cells.

45 (2022) For the L6 light chain chimeric gene, the Sall to <u>Lindlill</u> fragment of the mouse V_c module was pined to the human C_c module by the procedure outlined in FIGURE 18, forming plN02119. Replacement of the nee sequence with the <u>E, coil</u> got gene derived from pSV2-gpt resulted in pIN02120, which expressed L6 chimeric light chain and confers movechenolic acid resistance when transfected into mammalian coll.

[0203] The inclusion of both heavy and light chain chimeric genes in the same plasmid allows for the introduction to translected cells of a 1.1 gene ratio of heavy and light chain genes leading to a balanced gene dosage. This may improve expression and decrease manipulations of translected cells for optimal chimeric antibody expression. For this purpose, the DNA fragments derived from the chimeric heavy and light chain genes of pING2111 and pING2119 were combined into the expression plasmid pING2114 (FIGURE 19). This expression plasmid contains a selectable near marker and separate transcription units for each chimeric gene, each including a mouse heavy chain enhancer.

55 [0204] The modifications and V-C joint regions of the L6 chimeric genes are summarized in FIGURE 20.

- (6) Stable Transfection of Mouse Lymphoid Cells for the Production of Chimeric Antibody.
- [0205] Electroporation was used (Potter <u>et al. supra;</u> Toneguzzo <u>et al. Mol. Cell Blot.</u> 6.703 1986) for the introduction of L6 chimeric expression plasmid DNA into mouse Sp200 cells. The electroporation technique gave a transfection frequency of 1-10 x 10⁵ for the Sp2/0 cells.
- [0205] The two gene expression plasmid piNG2114 was linearized by digestion with AgIII restriction endonuclease and transfected into Sp202 cells, giving approximately fifty C418 resistant clones which were screened for human heavy and light chain synthesis. The levels of chimeric antibody chain synthesis from the two producers, D7 and SE3, are shown in Table 3. Chimeric L6 antibody was prepared by culturing the D7 transfectant cells for 24 hours at 21x16 cells will in 51 DMEM supplemented with HEPES buffer and penicillin and streptomycin. The supernatant was concentrated over an Amizon YMS0 membrane in 10 mMs doctiom phosphate buffer, pH8.0. The preparation was loaded over a DEAE-Callubose column, which separated the immunoglobulin into unbound and bound fractions. Samples from the DEAE-unbound, DEAE-bound and the pre-DEAE preparations (rem 1.5 ut of medium) was separately purified by affinity chromatography on a Protein-A Sepharose column, etking with 0.1 M sodium citrate, pH 3.5. The cluted antibody was neutralized and concentrated by Amison centricen filtration, in phosphate buffered salien. The yields for the perpearations were 12ug (DEAE unbound), 6ug (DEAE bound), and 9ug (pre-DEAE column). Western analysis of the antibody chains indicated that they were combined in an H₂L2 textmenr like native immunoglobulins.
- (7) A second purification for Chimeric L6 Antibody Secreted in Tissue Culture.

[0207]

25

30

- a. Sp2/0 plNG2114.D7 cells were grown in culture medium [DMEM (Gibco #320-1965), supplemented with 10% Fetal Bovine Serum (Hyclone #A-1111-D), 10mM HEPES, 1x Glutamine-Pen-Strep (Irvine Scientific #9316) to 1 x 10⁶ cell/ml.
- b. The cells were then centrifuged at 400xg and resuspended in serum-free culture medium at 2 x 10⁶ cell/ml for
- c. The medium was centrifuged at 4000 RPM in a JS-4.2 rotor (3000xg) for 15 min.
- d. 1.6 liter of supernatant was then filtered through a 0.45 micron filter and then concentrated over a YM30 (Amicon Corp.) filter to 25ml.
 - e. The conductance of the concentrated supernatant was adjusted to 5.7-5.6 mS/cm and the pH was adjusted to 8.0. f. The supernatant was centrifuged at 2000xg, 5 min., and then loaded onto a 40 ml DEAE column, which was
- preequilibrated with 10mM scollum phosphate, pH8.0.
 g. The flow through fraction was collected and loaded onto a 1ml protein A-Sepharose (Sigma) column preequilibrated with 10mM sodium phosphate, pH8.0.
 - h. The column was washed first with 6ml 10mM sodium phosphate buffer pH=8.0, followed by 8ml 0.1M sodium citrate pH=3.5, the by 6ml 0.1M citric acid (pH=2.2). Fractions of 0.5ml were collected in tubes containing 50ul 2M fris base (Sigma).
 - i. The bulk of the IgG was in the pH=3.5 elution and was pooled and concentrated over Certricon 30 (Amicon Corp.) to approximately .06ml.
 - j. The buffer was changed to PBS (10mM sodium phosphate pH=7.4, 0.15M NaCl) In Centricon 30 by repeated diluting with PBS and reconcentrating.
 - k. The IgG solution was then adjusted to 0.10ml and bovine serum albumin (Fraction V, U.S. Biochemicals) was added to 1.0% as a stabilizing reagent.
 - (8) Production and Purification of Chimeric L6 Antibody Secreted in Ascites Fluid.

[0208]

50

a. The ascites was first centrifuged a 2,000 xg for 10 min.

of 0.5 M NH₄OH, and 3 M sodium thiocyanate.

- b. The conductance of the supernatant was adjusted to 5.7-5.6 mS/cm and its pH adjusted to 8.0.
- c. Supernatant was then loaded onto a 40 ml DEAE-cellulose column pre-equilibrated with 10 mM Na₂PO₄H pH 8.0.
- d. The flow through from the DEAE column was collected and its pH was adjusted to 7.4, and then loaded onto a 1.0 ml goet anti-human IoG (H + L) sepharose column.
- e. The colump was washed first with 6 ml of 10 mM sodium phosphate, 0.5 M sodium chloride, followed by 8 ml
 - f. The sodium thiocyanate cluate was pooled and dialyzed against 2L PBS overnight.

[0209] The antibody can be further concentrated by steps i, and k, of the previous procedure.

TABLE 3

Culture Condition	FBS Sp2/0.D7		Sp2/0.3E3		
		<u>Kappa</u> b	<u>Gamma</u> ^e	Kappab	<u>Gamma</u> ^c
1. 20 ml, 2d, seed @ 2x10 ⁵ /ml	+	17	77	100	700
2. 200 ml, 2d, seed @ 2.5x10 ⁵ /ml	+	0.9	6	80	215
3. 200 ml, 1d, seed @ 2x106/ml	- 1	1.9	3.8	97	221
4. Balb/c ascites	- 1	5,160	19,170	ND	ND

FBS: Fetal Bovine Serum

10

15

- a Sp2/0 cells transfected by electroporation with pING2114(pL6HL)
- b ug/l measured by ELISA specific for human Kappa human Bence-Jones protein standard
- c ug/l measured by ELISA specific for human gamma human IgG standard.
- 20 (9) Studies Performed on the Chimeric L6 Antibody.

[0210] First, the samples were tested with a binding assay, in which cells of both an L6 antigen-negative cell line were incubated with standard mouse monoclonal antibody L6, chimeric L6 antibody deviced from the cell culture supernatants, and chimeric L6 antibody deviced from ascites (as previously described) followed by a second reagent, fluorescein-isothiocyanate (FITC)-conjugated goat antibodies to human (or mouse, for the standard) immunoclobulin.

[0211] Since the binding assay showed strong reactivity of the chimeric L6 on the L6 antigen positive cell line and total lack of reactivity on the negative cell line, the next step was to test for the ability of the chimeric L6 to inhibit the binding of mouse L6 to antigen positive cells; such inhibition assays are used routinely to establish the identity of two antibodies' recognition of antigen. These data are discussed below ("Inhibition of binding"). As part of these studies, a rough estimate of antibody avidity was made.

[0212] Finally, two aspects of antibody function were studied, the ability to mediate ADCC in the presence of human peripheral blood leakcytes, and the ability to kill Le positive tumor cells in the presence of human serum as a source of complement (see *Functional Assays* below).

56 [Q13] <u>Binding Assays</u>, Cells from a human colon carchroma line, 347, which had been previously shown to express approximately 5 x 10⁸ molecules of the L6 antigen at the cell surface, were used as targets. Cells from the T cell line HSB2 was used as a negative control, since they, according to previous testing, do not express detectable amounts of the L6 antigen. The target cells were first incubated for 30 min at 4°C with either the chirerie L6 or with mouse L6 standard, which had been purified from mouse sacties. This was followed by incubation with a second, FITC-labelled, reagent, which for the chireric antibody was goat-anti-human immunoglobulin, obtained from TAGO (gotto), and used at a dilution of 1:50. For the mouse standard, it was goat-anti-nouse immunoglobulin, also obtained from TAGO and used at a dilution of 1:50. Antibody binding to the cell surface was determined using a Coulter Model EPICC cell sorfer.

[0214] As shown in Table 4 and Table 4A, both the chimeric and the mouse standard L6 bound significantly, and to approximately the same extent, to the L6 positive 3341 line. They did not bind above background to the L6 negative

[0215] In view of the fact that the three different chimeric L6 samples presented in Table 4 behaved similarly in the binding assays, they were pooled for the inhibition studies presented below. The same inhibition studies were performed for chimeric L6 derived from ascites fluid presented in Table 4A.

[0216] Inhibition of Binding. As the next step was studied the extent to which graded doses of the chimeric L6 antibody, or the standard mouse L6, could inhibit the binding of an FITC-tabelled mouse L6 to the surface of antigen positive 3347 coton carcinoma cells.

[0217] Both the chimeric and mouse standard L6 inhibited the binding of the directly abelled L6 antibody, with the binding curves being parallel. The chimeric antibody was slightly less effective than the standard, as indicated by the results which showed that 3.4 ug/ml of the pooled chimeric L6 MAb, as compared to 2.0 ug/ml of the standard mouse L6 MAb was needed for 50% inhibition of the binding, and that 5.5 ug/ml of the chimeric L6 (derived from ascites) as compared to 2.7 ug/ml of the standard mouse L6 MAb was needed for 50% inhibition of binding.

[0218] As part of these studies, a rough estimate was made of antibody avidity. The avidity of the standard mouse

L6 had been previously determined to be approximately 4 x 108. The data indicated that there were no significant differences in avidity between the chimeric and the mouse L6.

[0219] <u>Functional Assays</u> A comparison was made between the ability of the chimeric L6 and standard mouse L6 to yes L6 antigen positive cells in the presence of human perhipheral blood leukocytes as a source of efector cells (madialing Antibody) Dependent Cellular Cytotoxcity, ADCC) or human serum as a source of complement (mediating Complement-Dependent Cytolysis, CDC).

[0220] As shown in Table 5 and Tables 5A-5D, the chimeric L6 was superior to the simultaneously tested sample of mouse L6 in causing ADCC, as measured by a 4 hr ⁵¹Cr release test.

[0221] Tables 6 and 6A-6B present the data from studies on complement-mediated target cell lysis. In this case, a high cytolytic activity was observed with both the mouse and the chimeric L6 antibodies.

Conclusions

50

[0222] The results presented above demonstrate a number of important, unexpacted qualities of the chimeric L6 monoclonal antibody of the invention. Firstly, the chimeric L6 antitibody binds to L6 antitigen positive turnor cells to approximately the same extent as the mouse L6 standard and with approximately the same avidity. This is significant for the following reasons: the L6 antibody defines (a) a surface carbodydrate antigen, and (b) a protein antigen of about 20,000 dattons, each of which is characteristic of non-small cell lung carcinoma (NSCLC) and certain other human carcinomas. Significantly, the L6 antibody does not bind detectably to normal cells such as fibroblests, endothelial

carcinomas. Significantly, the L6 antibody does not bind detectably to normal cells such as fibroblasts, endothelial cells, or epithelial cells in the major organs. Thus the chimeric L6 monoclonal antibody defines an antigen that is specific for carcinoma cells and not normal cells.

[0233] In addition to the ability of the chimeric L6 monoclonal antibodies of the present invention to bind specifically.

to malignant cells and localize furnors, the chimeric L6 exerts profound biological effects upon binding to its target, which make the chimeric antibody a prime candidate for tumor immunotherapy. The results presented herein demonstrate that chimeric L6 is capable of binding to tumor cells and upon binding falls the tumor cells, either by ADCC or CDC. Such tumor killing activity was demonstrated using concentrations of chimeric L6 antibody as low as 0.01 ug/nl (10ng/ml).

[3244] Although the prospect of attempting tumor therapy using monoclonal antibodies is attractive, with some partial tumor, regressions being reported, to date such monoclonal antibody through some met with limited success of (Houghton, February 1985, <u>Proc. Natl. Acad. Sci. B2</u>:1242-1246). The therapsutic afficacy of mouse monoclonal antibodies (which are the ones that have been tried so pria) appears to be too low four most practical purposes. The discovery of the profound biological activity of chimeric L6 coupled with its specificity for a carcinoma antigan makes the chimeric L6 antibody a choice therapsutic agent for the retentant of tumors <u>in yivo</u>, Moreover, because of the "human" properties which will make the chimeric L6 monoclonal antibodies more resistant to clearance <u>in yivo</u>, the chimeric L6 monoclonal antibodies more resistant to clearance <u>in yivo</u>, the chimeric L6 monoclonal antibodies more resistant to clearance to the "tumor of development of various immunoconjugates with drugs, toxins, immunomodulators, isotopes, etc., as well as for diagnostic purposes such as <u>in yivo</u> maging of tumors using appropriately labelled chimeric L6 antibodies. Such immunoconjugation techniques are known to those skilled in the art and can be used to modify the chimeric L6 antibody molecules of the present invention.

40 [0225] Two illustrative cell lines socreting chimeric L6 antibody were deposited prior to the filing date of this application at the ATCC, Rockville Maryland. These are transfected hybridoma C255 (corresponds to 3E3 cells, <u>supra</u>), ATCC HB 9240 and transfected hybridoma C256 (C7 cells, <u>supra</u>), ATCC HB 9241.

[0225] The present invention is not to be limited in scope by the cell lines deposited since the deposited embodiment is intended as a single liuration of one aspect of the invention and all cell lines which are functionally equivalent are the within the scope of the invention. Indeed, various modifications of the invention in addition to those shown in the art from the foreigning description and accompanying drawings are intended to fall within the scope of the appended claims.

TABLE 4

		IAL	700 7	
	Binding Assays of Chimeric		6 Monoclonal Antibody on an ative Cell Line.	L6 Antigen Positive and L6
			Binding Ratio For* H3347 Cells (L6 +)	
	Antibody	Batch	GAM	GAH
,	Standard L6		56.6	4.2
´	Chimeric L6	а	1.3	110.3
	1	ь	1.3	110.3
-		c	1.3	110.3
H			Binding Ratio For* HSB-2 Cells (L6 -)	
			GAM	GAH
.	Standard L6		1.1	1.1
	Chimeric L6	· a	1.0	1.0
- 1		b	1.0	1.1
- 1		c	1.0	1.1

^{*}All assays were conducted using an antibody concentration of 10 ug/ml. The binding ratio is the number of times brighter a test sample is than a control sample treated with GAM (FTIC conjugated post anti-human) alone. A ratio of 1 means that the test sample is just as bright as the control, a ratio of 2 means the test sample is wise as bright as the control.

25

TABLE 4A

	Antibody Concentration	Binding Ratio For* H3347 Cells (L6 +)	
Antibody	(ug/ml)	GAM	GAH
Standard L6	30	38	. 4
	10	49	4
	3	40	3
Chimeric L6	30	2	108
(Ascites)	10	2	108
	3	1	42
Chimeric L6	30	1	105
(Cell culture)	10	1	86
	3	1	44
		Binding Ratio For	** HSB-2 Cells (L6 -)
		GAM	GAH
Standard L6	10		

The binding ratio is the number of times brighter a test sample is than a control eample treated with GAM (FITC conjugated gost anti-human) alone. A ratio of 1 means that the test sample is just as bright as the control, a ratio of 2 means the test sample is twice as bright as the control, etc.

TABLE 4A (continued)

Binding Assays Of Ch	imeric L6 Antibody and Mouse Monocl Negative Ce		igen Positive and L6 Ant
	Antibody Concentration	H3347 Cells (L6 +)	
		Binding Ratio For** HSB-2 Cells (L6 -)	
		GAM	GAH
Chimeric L6	10	1	1
(Cell Culture)			

10

35

TABLE 5

	ADCC of Chimeric L6 (Mouse) L6 Antibodies On Colon Carcinoma Cell Line 3347.			
20	Antibody	Antibody Concentration (ug/ml)	PBL per Target Cell	% Cytolysis*
	Chimeric L6	10	100	64
- 1		5	100	70
25		10	0	2
	Standard L6	10	100	24
		5	100	17
		10	0	2
30	None	0	100	1

The larget cells had been labelled with ⁵¹Cr and were exposed for 4 hours to a combination of MAb and human peripheral blood feukocytes (PBL), and the release of ⁵¹Cr was measured subsequently. The release of ⁵¹Cr (after corrections of values for epontaneous release from untreated cells) is a measure of the percent cytolist.

TABLE 5A

	TABL	E SA		
ADCC of Chimeric L6 and Standard (Mouse) L6 Antibodies On Colon Carcinoma Cell Line 3347.				
Antibody	Antibody Concentration (ug/ml)	PBL per Target Cell	% Cytolysis*	
Chimeric L6 (Ascites)	20	100	80	
	10	100	74	
	5	100	71	
i	2.5	100	71	
	20	0	0	
Chimeric L6 (Cell Culture)	10	100	84	
	5	100	74	
	2.5	100	67	
	10	0	3	
Standard L6	20	100	32	
	10	100	26	
	20	0	0	

^{The turget cells had been labelled with ⁵1 Cr and were exposed for 4 hours to a combination of MASs and human peripheral blood suckerytes (PBL), and the refease of ⁵1 Cr was measured subsequently. The release of ⁵1 Cr (after corrections of values for spentaneous release from unrested cells is a measure of the percent cytobia.}

[&]quot;The binding ratio is the number of times brighter a test sample is than a control sample treated with GAM (FITC conjugated goat anti-human) alone.

A ratio of 1 means that the test sample is just as bright as the control, a ratio of 2 means the test sample is twice as bright as the control, etc.

TABLE 5B

Antibody	Antibody Concentration (ug/ml)	PBL per Target Cell	% Cytolysis*
Chimeric L6 (Ascites)	5	100	84
	2.5	100	78
	1.25	100	85
	0.63	100	81
	0.31	100	80
	0.16	100	71
	0.08	100	65
	5	0	0
Standard L6	5	100	32
	5	0	0
None	0	100	19

The target cells had been labelled with ⁵1Cr and were exposed for 4 hours to a combination of MAB and human periphenal blood isviscoyles (PBL), and the release of ⁵1Cr was measured subsequently. The release of ⁵1Cr (after corrections of values for spontaneous release from untrested cells) is a measure of the percent rejoicits.

TABLE SC

Antibody	% Cytolysis*		
Antibody	Antibody Concentration (ug/ml)	PBL per Targtet Cell	76 Cytolysis
Chimeric L6	10	100	35
(Ascites)	1	100	31
	0.1	100	27
	0.01	100	15
	0.001	100	13
	0.0001	0	15
Standard L6	, 10	100	9
	1	100	15
None	0	100	9
Chimeric L6	10	10	19
(Ascites)	1	10	15
	0.1	10	11
	0.01	10	13
	0.001	10	22
	0.0001	10	11
Standard L6	10	10	7
	1	10	6

^{*}The targel cells had been labelled with 5¹Cr and were exposed for 4 hours to a combination of IMAb and Human peripheral blood loukocytes (PBL), and the release of 5¹Cr was measured subsequently. The release of 5¹Cr (after corrections of values for Spontaneous release from untreated cells) is a measure of the percent of loyless.

TABLE 5C (continued)

ADOC of Chimeric	ADOC of Chimeric L6 and Standard (Mouse) L6 Antibodies On Lung Carcinoma Cell Line H2669.					
Antibody	Antibody Concentration (ug/ml)	PBL per Targtet Cell	% Cytolysis*			
None	0	10	8			
Chimeric L6 (Ascites)	10	0	4			
Standard L6	10	0	9			

^{*}The target cells had been labelled with ⁵¹Cr and were exposed for 4 hours to a combination of IMAb and Human pertipheral blood butkcoytes (PBL), and the release of ⁵¹Cr was measured subsequently. The release of ⁵¹Cr (after corrections of values for Spontaneous release from unterested cells) is a measure of the personn dyshybat.

TABLE 5D

15

35

40

50

55

ADCC of Chimeric L6 and Standard (Mouse) L6 Antibodies On Colon Carcinoma Cell Line H3347.					
Antibody	Antibody Concentration (ug/ml)	PBL per Target Cell	% Cytolysis*		
Chimeric L6 (Ascites)	10	100	62		
	1 1	100	66		
	0.1	100	69		
	0.01	100	26		
	0.001	100	8		
	0.0001	0	3		
	10	0	0		
Standard L6	10	100	19		
	1 1	100	24		
		0	0		
None	0	100	8		

^{*} The target celle had been labelled with ⁵¹Cr and were exposed for 4 hours to a combination of MAb and Human peripheral blood leukocytes (PBL), and the release of ⁵¹Cr (after corrections of values for spontaneous release from untreated cells) is a measure of the percent cytolysis.

TABLE 6

of complement.						
ŀ	line 3347, as measured by a 4-hr 51 Cr-release assay. Human serum from a healthy subject was used as the source					
-[Complement-dependent cytotoxic effect of chimeric and standard (mouse) L6 on colon carcinoma cells from					

	Antibody	Human complement	% Cytolysis
	L6 Standard 10 ug/ml	Yes	90
45	L6 chimeric 10 ug/ml	Yes	89
	L6 Standard 10 ug/ml	No	0
	L6 chimeric 10 ug/ml	No	1

TABLE 6A

	INDEE ON		
Complement Dependent Cyto	toxic Effect of Chimeric L6 and Sta Cell Line 3347		s on Colon Carcin
Antibody	Antibody Concentration (ug/ ml)	PBL per Target Cell	% Cytolysis
Chimeric L6 (Ascites)	20	+	29
	10	+	23
	5	+	18
	2.5	+	8
	20	Inactivated	0
	10	0	0
Chimeric L6 (Cell Culture))	20	+	29
	5	+	26
	2.5	+	18
	20	+	4
	10	0	4
Standard L6	20	+	55
	10	+	37
	20	Inactivated	0
	20	0	1
			0

^{*} Complement mediated cytolysis was measured by a 4 hour ⁵¹Cr-release assay. Human serum from a healthy subject was used as the source of complement.

TABLE 6B

	1 12 1 2 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1	DDI To C-II	% Cytolysis*
Antibody	Antibody Concentration (ug/ml)	PBL per Target Cell	% Cytolysis
Chimeric L6 (Ascites)	10	+	209
	5	+	155
	2.5	+	166
	1.25	+	114
	0.6	+	63
	0.3	+	17
	10	0	0
Standard L6	10	+	96
	5	+	83
	2.5	+	48
	1.25	+	18
	0.6	+	7
	0.3	+	4
	10	0	2

^{*} Complement mediated cytolysis was measured by a 4 hour ⁵¹Cr-release assay. Human serum from healthy subject was used as the source of complement.

EXAMPLE IV: A Human-Mouse Chimeric Antibody with Specificity for Human B-Cell Antigen

[0227] The ZHT mouse monoclonal antibody (gamma ZhK) recognizes a human B-cell surface antigen, Bp35 (Clark, E.A., et al., Proc. Nat. Acad. Sci. USA 82:1766 (1985)). The Bp35 molecule plays a role in B-cell activation. mRNA was prepared from the 2H7 cell line. Two CDNA libraries were generated - one using the heavy chain UIG-H primer and the other, oligo(dT). One V_H clone, pH2-11, was isolated upon screening with the same UIG-H oligonucleotide. To isolate the light chain clone, a mouse largue-specific DNA fragment was used to screen the oligo(dT) library. Candidate clones were further screened with a mouse lags expeciences. One V_K clone, pL2-12, was thus isolated. The light chain UIG-K was then used to encineer a restriction enzyme site in the J goolon.

[0228] The two cDNA clones were also modified at the 5' end to remove the artificial otigo dfC] sequence. In pH2-11 this was carried out by using the restriction enzyme N₂₀₀ which out so ne nucleotide residue 5' of the ATG initiator codon. In pL2-12 this was achieved by an oligonucleotide in <u>vitto</u> mutagenesis using a 22-mer container a Sall site. [0229] The DNA sequences of these two clones are shown in Figures 21, 22. To construct the chimeric heavy chain plasmid, the V_x module was joined to the human C gamma 1 module (pGML60) at the J_x <u>Hindli</u> site, and the chimeric light chain the V_x module was joined to the human C gamma 1 module (pGML60) at the J_x <u>Hindli</u> site, and the chimeric light chain the V_x module was joined to the human C_x module (pGML60) at the J_x <u>Hindli</u> site. The expression vector sequences were derived from pinXc2012-neas well as pinXc2016-gpt. The constructed plasmids are pinXc2010 (C_y gamma 1-neo), pinXc2106 (V_xC_x-neo), pinXc2101 (V_xC_x-rgpt), pinXc2101 and pinXc2106 were also used to generate plasmids containing both genes. They are pH12-11 and pH12-26. In addition, pinXc2106 and pinXc2014 were combined to a two light chain plasmid, p12-25, to compensate for the power (compared to heavy chain) steady state accumulation of light chain protein in transfected cells. (See Fig. 23.) Fig. 24 shows the changes made to the variable region sequences during the construction.

[0230] The plasmid, phtl.2-11, was linearized by Agill; and the DNA was used to transfect Sp20 cells by electropation. Transformants were selected in G418-DMEM. One transformant, 1C9, produces 8.3 ng/m chimeric kappa and 33-72 ng/ml chimeric gamma 1 protein as assayed by ELISA. Southern analysis of 1C9 DNA showed that there is one copy of the plasmid integrated in Sp20 genome.

EXAMPLE V: Secretion of a Functional Chimeric Antibody from Yeast

(1) Fusion of mature chimeric L6 light chain and heavy chain genes to the yeast invertase signal sequence and shortened phosphoglycerate kinase (PGK promoter).

[0231] Yeast cells are capable of recognizing mammalian secretion signal sequences and of directing secretion of mammalian proteins (Hitzman et al., supra). There is, however, evidence which suggests that certain native yeast signal sequences are more effective than mammalian signal sequences at directing secretion of some mammalian proteins from yeast (Smith et al., Science 229:1219 (1985)). One example is the signal sequence for the yeast invertase gene. To improve the efficiency of light and heavy chain secretion, the mature light chain and heavy chain sequences were fused to the yeast invertase signal sequence and placed under transcriptional control of the shortened PGK promoter (U.S. Patent Application 797,477) using the strategies outlined in Figures 25 and 26, respectively. An important element of these constructions is the use of in vitro mutagenesis to introduce a restriction site at the signal sequence processing site for both the invertase signal sequence (see U.S. Patent Application 797,477) and the light and heavy chain genes. These restriction sites are positioned such that a blunt-ended ligation of restriction enzyme-digested, T-4 DNA polymerase-treated DNA results in in-phase translational fusions of the 5' end of the mature immunoglobulin chains with the 3' end of the yeast invertase signal sequence. Such genes, when expressed in a yeast cell, may direct the synthesis, processing, and secretion of chimeric light and heavy chains with the same primary peptide sequence as chimeric light and heavy chains secreted from transfected mouse Sp2/0 cells. The DNA sequences of the mutagenesis primers used for light and heavy chain genes as well as the corresponding unmutagenized sequences are shown in Figures 25B and 26B, respectively. Using this approach, the L6 chimeric light and heavy chains were fused to the yeast invertase signal sequence and shortened PGK promoter, resulting in plasmids pING1407-7 and pING1415 (Figures 25C and 26C).

(2) Removal of non-yeast 3' untranslated DNA.

50

[0232] Recent studies on expression of hepatitis B surface antigen in yeast demonstrated that removal of non-yeast 3 and 5 untranslated sequences can result in increased levels of heterologous gene expression in yeast (Rinsking it al. Septe 46: 135 (1989)). The light chain gene sequence of chimeric L6 antibody in pING1407-7 (Figure 25C) contains approximately 200 bp of 30 untranslated DNA (clickwed by 70 bp of pdy A and 20 bp of poly Sequences. An initial treatment of the chimeric L6 light chain DNA with the double-stranded exonuclease Ball 31, removed the poly A and poly Sequences and all but 90 bp of 30 untranslated DNA, generating the plasmid pING2121b (Figure 27). A restriction

Iragment from pINQ2121b containing only C_s was choned into a derivative of pBR322, generating pING1419 (Figure 27). A second <u>Bal</u>31 digestion was next used to remove all but 13 bp of non-yeast 3' untranslated DNA generating the plasmid, pING1431 (Figure 27). The chimeric L5 heavy chain gene in pING1415 (Figure 25) also contains extensive 3' untranslated sequence which includes 80 bp of poly A. All but 11 bp of the 3' untranslated DNA were removed using the strategy shown in Figure 28, generating the plasmid pING1429.

[023] Site-directed in <u>vitro</u> multagenesis can introduce, at a low frequency, unwanted base pair changes in regions of the DNA outside of the area being multagenized. To ensure that such mutations were not present in the iligin and neavy chain sequences which had been cloned into M13 and subjected to site-directed multagenesis, we constructed light and heavy chain genes lused to the invertase signal sequence and the shortened PGK promoter which consisted of coding sequences that were either confirmed by DNA sequence analysis or proven to be functional by virtue of their expression in transfected mouse Sp200 cells to produce functional chimeric L6 antibody. The plasmids, plots 1399 (light chain, Figure 27) and pING1439 (light chain, Figure 27) and pING1439 (light seponstated by these constructions).

(3) Construction of yeast expression plasmids containing chimeric L6 light and heavy chain genes from pING1439 and pING1436, respectively, fused to the PGK polyadenylation signal.

[0234] In order for yeast to produce an intact functional antibody molecule, a balanced synthesis of both light and heavy chain protein within the host cell is preferred. One approach is to place the light and heavy chain genes on separate expression vectors each containing a different selective marker. A yeast strain defective in the selective 20 markers found on the plasmids can then be either simultaneously or sequentially transformed with these plasmids. [0235] The chimeric L6 light and heavy chain genes from pING1439 (Figure 27) and pING1436 (Figure 28) were cloned as BqIII-Xhol and BamHI-Xhol fragments, respectively, in two different medium copy number (about 20 copies/ cell) expression vectors (yeast-E. coli shuttle). One of these, pING804CVS, contains the complete yeast 2-micron circle, the PGK transcription termination and polyadenylation signals, and the leu2 gene as the selective marker. The 25 other vector, pING1150, contains the yeast origin of replication, oriY, a cis-acting sequence (REP3) from the yeast endogenous 2-micron plasmid, the PGK transcription termination and polyadenylation signals, and the ura3 gene as the selective marker. Both plasmids also contain the β -lactamase gene (bla) for ampicillin resistance and the bacterial origin of replication (oriB) from pBR322 for selection and amplification in bacteria. Four plasmids resulted from these constructions; pING1441--light chain, leu2 and pING1443--light chain, ura3 (Figure 29); pING1440--heavy chain, leu2 and pING1442--heavy chain, ura3 (Figure 30).

(4) Secretion of chimeric L6 antibody from transformed yeast cells.

55

[0236] Two separate transformation experiments were performed in an attempt to obtain both light and heavy chain synthesis in yeast cells. Four µg each of pING1440 and pING1443, and separately of pING1442 and pING1441 were cotransformed into Saccharomyces cerevisiae strains BB331C (MATa, ura3, leu2) by selecting for growth on SD agar (2% glucose, 0.67% yeast-nitrogen base, 2% agar). Ura+ Leu+ transformants appeared at 2-3 days of incubation at 30°C. Approximately 100 transformants were obtained for pING1440 plus pING1443; only 15 transformants were obtained for pING1442 plus pING1441. Ten colonies were inoculated from each plate into 5 ml SD broth supplemented with 50 mM sodium succinate, pH 5.5, and grown for 65 hours at 30°C. The cells were removed by centrifugation and the culture supernatants analyzed by ELISA for the levels of light chain and heavy chain and for the degree of association of the secreted light and heavy chains. The latter was assessed using a goat anti-human kappa antiserum to coat the micro-titer wells and a peroxidase-labeled goat anti-human gamma antiserum to detect the level of heavy chain bound to the anti-kappa coat. The results of these assays (Table 7) revealed that all of the culture supernatants from the cells transformed with pING1440 (heavy chain, leu2) plus pING1443 (light chain, ura3) contained a disproportionately high level of light chain protein relative to the levels of heavy chain protein, and no evidence (at least as determined by ELISA) of assembled light and heavy chains. On the other hand, the supernatants from the cells transformed with pING1442 (heavy chain, ura3) + pING1441 (light chain, leu2) contained a more balanced production of light and heavy chain proteins, and eight of ten isolates appeared to contain some assembled light and heavy chains as determined by ELISA. Two of these isolates, No. 1 and No. 5, produced a significant proportion of assembled light and heavy chain.

TABLE 7

			I ADLE /			
	LEVELS OF SECRET	LEVELS OF SECRETED CHIMERIC L6 LIGHT AND HEAVY CHAIN BY YEAST TRANSFORMANTS ^a				
	Plasmids ^b	Isolate No.	Kappac	<u>Gamma</u> ^d	Kappa/Gamma*	
5	pING1440+ pING1443	1	284	39	0	
		2	324	33	0	
		3	473	52	0	
	+	4	387	40	0	
10		5	316	34	0	
		6	188	28	0	
		7	381	45	0	
		8	455	45	0	
15		9	380	26	0	
15		10	579	32	0	
	pING1441 + pING1442	1	128	79	35	
		2	150	30	1	
		3	124	29	0 .	
20		4	185	55	5	
		5	114	52	35	
		6 '	139	23	5	
		7	149	34	5	
25		8	245	57	12	
		9	202	26	11	
		10	157	19	7	

a. S. cerevisiae strain BB331C (MATa, leu2, ura3) transformed to Ura+ Leu+ with plasmids carrying ura3 and leu2 with light or heavy chains.

30

35

[0237] Further analysis was performed to determine if this association was the result of the synthesis of an H₂L₂ size protein. The culture supernatants from isolates Nos. 1 and 5, as well as from isolate No. 8, which contained a much lower level of apparent light and heavy chain association, were concentrated by ultrafilitation on a Centricton 30 filter (Amicon Corp.). The concentrated supernatants were run on a 7% polyacrylamide gel under non-reducing conditions, blotted to nitrocellulose, and probed with goat anti-human kappa antiserum followed by peroxidase-ided rabbit anti-goat antiserum. The concentrated supernatants from isolates No. 1 and 5, but not from No. 8, contained a single immunoreactive band which comigrated with his purified chimeric L6 antibody from transfected Sp20 cells. These results suggested that isolates No. 1 and 5 were synthesizing and secreting assembled L6 chimeric antibody.

(5) Purification of chimeric L6 antibody from yeast culture supernatant.

[0238] Inorder to further characterize the hgl_scize protein secreted by the yeast and determine if this was assembled to chimeric antibody, a sufficient quantity of yeast-produced material was purified to allow the performance of various binding and functional assays. The pING 1442 + 1441 transformant isolate No. 5 was grown for 58 hours at 30°C in a 10-liker fementor using a synthetic medium (Table 8). The cells were initially grown in 3 liters of the column A medium until the glucose level fell below 1 gft_at which time they were fed with a total volume of 2.5 t. of medium from column 8. Glucose levels were maintained at 1.5 gft_during the remaining course of the fermentation. The cells were removed by centrifugation and the culture supernatural was analyzed by ELISA for the presence of light end heavy chain proteins and for association of the heavy and light chains. The supernatural two processes are consistent of the heavy and light chains. The supernatural two processes are concentrated by stratification over a DC. 10 unit (Amicon Corp.), filtered through 0.45 micron litter and concentrated over a YM50 little (Amicon Corp.) to 250 ml. The concentrated adjusted to pl 1-2 with KOH, brought

pING1440 = heavy chain + <u>leu</u>2;

pING1443 = light chain + ura3;

pING1442 = heavy chain + ura3; pING1441 = light chain + leu2.

pind 1441 # light chain + leuz.

c. ng/ml measured by ELISA specific for human kappa with human Bence Jones protein as standard.

d. ng/ml measured by ELISA specific for human gamma with human as IgG standard.

e. ng/ml measured by ELISA using anti-human kappa as coating antibody and anti-human gamma as second antibody with human igG standard.

to 500 ml with PBS (10 mM sodium phosphate, pH 7.4, 150 mM sodium chloride) and loaded on a 1 ml protein A-Sepharose (Sigma) column, pre-equilibrated with PBS. The couldnm was washed first with 20 ml PBS, followed by 10 ml 0.1 M sodium citrate, pH 3.5, then by 10 ml 0.1 M citric acid pH = 2.2. The pH 3.5 and 2.2 eluates were each collected in a tube containing in ml 2 M Tris base (Sigma). The bulk of the light and heavy chain immunoreactive proteins were in the pH 3.5 eluate which was next concentrated over a Centricon 30 (Amicon Corp.) to a final volume of 106 ul. Analysis of this protein on non-reducing polyacrylamide gels using coomassis blue staining and immunoblotting with anti-human kappa antiserum (Sigma) to visuatize the proteins revealed an H₂L-pize, 155 (kidations, protein band. This protein was purified away from other proteins by HPLC using an AB, 5-micron column equilibrated with buffer A (10 mM KPQ₄, pH 6.8). After clading the sample on the column, the column was washed with buffer A for 10 minutes (flow rate = 1 ml/minute) and subjected to a linear gradient of 0% to 50% buffer B (250 mM KPQ₄, pH 6.8) over 50 minutes at 1 ml/minute.

TABLES

	IA.	BLE 8		
MEDIUM USED FOR YEAST FERMENTATION TO PRODUCE SECRETED L6 CHIMERIC ANTIBODY*				
Ingredients		Ab	Bc	
1.	Cerelose (Glucose)	119 g/l	538 g/l	
2.	(NH ₄) ₂ SO ₄	13.9 g/l	83.3 g/l	
3.	Thiamine HCL	0.011 g/l	0.05 g/l	
4.	Biotin	0.00011 g/I	0.005 g/l	
5.	Pantothenic acid	0.002 g/l	0.009 g/l	
6.	Inositol	0.194 g/l	0.875 g/l	
7.	H ₃ PO₄	5.67 ml/l	25.5 ml/l	
В.	KH ₂ PO ₄	5.78 g/l	26.0 g/l	
9.	MgSO ₄ .7H ₂ O	3.33 g/l	15.2 g/l	
10.	CaCl ₂ .2H ₂ O	0.33 g/l	1.5 g/l	
11.	FeSO ₄ .7H ₂ O	0.072 g/l	0.34 g/l	
12.	ZnSO ₄ .7H ₂ O	0.022 g/l	0.104 g/l	
13.	MnCl ₂ .4H ₂ O	0.0039 g/l	0.018 g/l	
14.	CuSO ₄ .5H ₂ O	0.0067 g/l	0.031 g/l	
15.	Conc. HoSO4	0.0056 mVI	0.026 mVI	

- a. Fermentation was performed as described in text.
- b. Constituents of Initial 9-liter batch.

35

45

c, Constituents of 2.5-liter feed batch.

[0239] The bulk of the protein resolved into a single large broad peak between 20 and 50 minutes as determined by absorbance at 280 nm. A second smaller peak was observed at 52-56 minutes, which corresponded to the normal outlon position for chimeric L6 artibody from translected Sp2/O cells. ELISA analysis of the column fractions revealed a major heavy + light chain cross-reactive peak corresponding to the U.A absorbance peak at 52-56 minutes. Analysis of the 52-56 minute fractions on non-reducing SDS polyacytamide gels using coormassis blue staining and immuno-blotting revealed an essentially pure protein which co-migrated with L6 chimeric antibody purified from transfected Sp2/O cells.

(6) Studies performed on the chimeric L6 antibody secreted by yeast.

[0240] The purified yeast-derived antibody was assessed for function in several ways. First, the purified antibody was tested for its ability to bind directly to an L6 antigen-positive cell line. Second, the antibody was tested for its ability to inhibit binding of mouse L6 antibody to antigen-positive cells. Finally, the purified antibody was tested for two aspects of antibody function—the ability to mediate ADCC in the presence of human peripheral blood leukocytes and the ability to kill L6 positive turnor cells in the presence of human complement.

[0241] <u>Direct Binding Assay.</u> Cells from a human colon carcinoma line, 3347, which expresses approximately 5 x 105 molecules of the L6 antigen per cell on the cell surface, were used as targets. Cells from the T cell line, 151, were used as a negative control since they, according to previous testing, do not express detectable amounts of the L6 antigen. The target cells were first incubated for 30 min at 4°C with either the \$p2/C cell- or yeast-derived chimeric L6 artibody or with mouse L6 antibody standard purified from mouse ascites. This was followed by incubation with FITC-labeled goat-anti-human immunoglobulin for the chimeric antibodies or with FITC-labeled goat-anti-mouse immu-

nogbbulin for the mouse standard. Both labeled antibodies were obtained from TAGO (Burlingame, Olar Jan used at a dilution of 1:50. Antibody bindings and used at a dilution of 1:50. Antibody bindings and used at a dilution of 1:50. Antibody bindings and the property of the property of

to approxir

(2443) Inhibition of Binding. As the next step, the yeast chimeric L6 antibody and the Sp2/O cell-derived chimeric L6 antibody were tested for their ability to inhibit the binding of an FITC-labeled mouse L6 antibody to the surface of antition-ossitive 3347 colon carcinoma cells.

[0244] Both the yeast-derived and Sp2/O-derived chimeric L6 antibodies inhibited the binding of labeled mouse L6 of antibody and the binding curves were parallel. Eased on the results of these studies, a rough estimate was made of antibody avidily. The avidity of the Sp2/O cell-derived chimeric L6 had been previously determined to be approximately 4 x 10⁸. The data indicated that there were no significant differences between the avidities of yeast-derived chimeric L6 and Sp2/O cell-derived chimeric L6 antipen.

[0245] <u>Functional Assays</u>. A comparison was made between the ability of the yeast-derived chimeric LS, 6p2/C cell-derived chimeric LB and standard mouse L6 antibodies to type L6 antigen-positive cells in the presence of human peripheral blood leukocytes as a source of effector cells mediating Antibody Dependent Cellular Cytotoxicity (ADCC). As shown in Table 10, the chimeric L6 from yeast was slightly better than 5p2/C-cell-derived chimeric L6 and as previously observed, both were superior to the standard mouse L6 in causing ADCC, as measured by a four-hour ⁵¹Cr release to the control of the standard mouse L6 in causing ADCC, as measured by a four-hour ⁵¹Cr release to the control of the standard mouse L6 in causing ADCC, as measured by a four-hour ⁵¹Cr release to the control of the standard mouse L6 in causing ADCC, as measured by a four-hour ⁵¹Cr release to the control of the standard mouse L6 in causing ADCC, as measured by a four-hour ⁵¹Cr release to the control of the standard mouse L6 in causing ADCC, as measured by a four-hour ⁵¹Cr release to the control of the standard mouse L6 in causing ADCC, as measured by a four-hour ⁵¹Cr release to the control of the standard mouse L6 in causing ADCC, as measured by a four-hour ⁵¹Cr release to the control of the standard mouse L6 in causing ADCC, as measured by a four-hour ⁵¹Cr release to the control of the standard mouse L6 in causing ADCC as measured by a four-hour ⁵¹Cr release to the control of the standard mouse L6 in causing ADCC as measured by a four-hour ⁵¹Cr release to the control of the standard mouse L6 in causing ADCC as measured by a four-hour ⁵¹Cr release to the control of th

[0245] A comparison was next made between the yeast-derived chimeric L6, Sp2/O cell-derived chimeric L5 and standard mouse L6 antibodies for their abilities to lyas L6 antigen-positive cells by complement-dependent cytoplysis (CDC) when human serum was used as the source of complement. The results of this comparison (Table 11) demonstrated that while both the Sp2/O-cell-derived chimeric L6 and standard mouse L6 antibodies exhibited high cytolytic activity, the yeast-derived L6 antibody failed to cause any cytolysis even at the highest antibody concentration. These results were unexpected and demonstrated that the yeast-derived antibody has new and unique properties.

(7) Conclusions

[0247] A process is disclosed by which yeast can be genetically engineered to secrete functional antibodies. The yeast-derived chimeric antibody in this example binds to the appropriate target antipen with approximately the same avidity as the chimeric antibody produced by lymphoid (sp20) cells. The yeast-derived antibody also displays elimits ADCC activity as does \$920-derived antibody. Unlike the \$920 cell-derived antibody antibody, also displays elimits and the production of a variety of monocional antibodies and chimeric antibodies can be applicable for the production of a variety of monocional antibodies and chimeric antibodies antibodies and eliminar antibodies and eliminary antibodies antib

TABLE 9

BINDING ASSAYS OF CHIMERIC L6 ANTIBODY PRODUCED BY YEAST OR MOUSE Sp2/O CELLS ON AN LE ANTIGEN-POSITIVE AND AN L6 ANTIGEN-NEGATIVE CELL LINE		
	Binding Ratio ^b for:	
Antibody ^a	H3347 Cells (L6 +)	T51 Cells (L6-)
Standard Mouse L6	95	1.0
Sp2/O Chimeric L6	116	1.0
Yeast Chimeric L6	116	1.0

a. All antibodies were used at a concentration of 10 µg/ml.

b. The binding ratio is the number of times brighter a test sample is than a control sample treated with FITC-conjugated second antibody. Goat antimous antibody was used as the second antibody for standard mouse L6 monoclonal antibody. Goal anti-human antibody was used as the second antibody or years and \$92.00 chimeric L6 antibody.

TABLE 10

ADCC OF CHIMERIC L6 ANTIBODY DERIVED FROM YEAST OR SP2/O CELLS AND STANDARD (MOUSE) L6 ANTIBODY ON COLON CARCINOMA CELL LINE 3347				
Antibody	Antibody Concentration (µg/ml)	% Cytolysis*		
Standard mouse L6	5.0	42		
	1.0	48		
Sp2/O Chimeric L6	1.0	96		
	01	71		

Antibody	Antibody Concentration (µg/ml)	% Cytolysis*
Standard mouse L6	5.0	42
	1.0	48
Sp2/O Chimeric L6	1.0	96
	0.1	71
	0.01	54
	0.001	37
Yeast Chimeric L6	1.0	114
	0.1	108
	0.01	76
	0.001	60
None	0	23

^{*} The target cells had been labeled with ⁵¹Cr and were exposed for four hours to a combination of MAb and human peripheral blood laukocytes at 100 per larget cell, and the release of ⁵¹Cr was measured subsequently. The release of ⁵¹Cr (after corrections of values for spontaneous release from untreated cells) is a measure of the percent cytolysis.

TABLE 11

	HUMAN COMPLEMENT-DEPENDENT CYTOTOXIC EFFECTS OF CHIMERIC L6 ANTIBODY PRODUCED BY YEAST OR MOUSE Sp2/O CELLS ON COLON CARCINOMA CELL LINE 3347				
30	Antibody	Antibody Concentration (µg/ml)	Complement ^a (+ or -)	Percent Cytolysis	
	Standard mouse L6	5	+	122	
		1 1	+	53	
		5	-	1	
35	Sp2/O Chimeric L6	5	+	73	
		1 1	+	22	
		0.1	+	. 5	
		5	-	2	
40	Yeast Chimeric L6	5	.+	3	
		1	+	2	
		0.1	+	4	
		5 ·	· ·	2	

a Human serum from a healthy subject was used as the source of complement.

b Complement-mediated cytolysis was measured by a four-hour ⁵¹C-release assay.

10

15

20

25

[0248] The Fab portion of IgG consists of a single light chain molecule coupled by a disulfide bridge to a single truncated heavy chain molecule consisting of the variable region and CH1 (Figure 31). This heavy chain fragment is known as Fd. Fabs are potentially useful for a variety of therapeutic and diagnostic procedures. In addition, they are amenable to production by microbial fermentation.

[0249] The usual method for production of Fab involves the digestion of intact IgG with papain (see Figure 31) followed by purification of the Fab away from the Fc fragments generated in the digest. While this procedure is relatively straightforward and can result in high yields of Fab, it is somewhat time-consuming in that it first requires the production and purification of whole antibody followed by generation and, finally, purification of Fab. Furthermore, one-third of the whole antibody molecule-the Fc portion (Figure 31)--is not utilized.

[0250] The recent advances in gene cloning and site-specific mutagenesis technology make possible a more direct

EXAMPLE VI: Secretion of Functional Chimeric Fab from Yeast

and simple attendance and simple attendance and simple attendance is approach, a stop codon is introduced in the heavy chair gain early and simple attendance attendance and simple attendance and si

(1) Introduction of a stop codon in the hinge region of L6 chimeric heavy chain.

[0251] The strategy for introduction of a stop codon into the hinge region of L6 chimeric heavy chain is outlined in Figure 32A. The location of the stop codon within the hinge region and the DNA sequence of the mutagenesis primer are shown in Figure 32B. The stop codon placement corresponds to amino acid 226 in Figure 31. This procedure generated the plasmid pING1402 containing an Fd gene which codes for a protein consisting of 228 amino acids and extends six amino acids and extends six amino acids also introduced a unique <u>Bcl</u>l site at the stop codon which can be readily utilized for subsequent manipulations of the 3° and of Fd. These include, but are not necessaryly limited to, emoval of heavy chain 3° untranslated DNA as well as the engineering of various types of modifications of Fd including the addition of coding sequences for specific amino acids and the production of fusion proteins.

(2) Fusion of the mature Fd gene to yeast invertase signal sequence and shortened PGK promoter,

²⁰ [0252] The strategy for fusion of the Fd gene to the yeast invertase signal sequence is outlined in Figure 33. This approach made use of the prior construction of the yeast invertase signal sequence-mature L6 heavy chain fusion (Figure 26) and utilized a unique Agal site in the J region of the chimeric L6 heavy chain to replace the constant region in plNG1415 consisting of C₁1, C₁2, and C₂3 with the constant region from plNG1412 containing the stop codon in the hinge region. This procedure generated the plasmid, plNG1418.

(3) Removal of non-yeast 3' untranslated DNA.

[0253] The introduction of a unique <u>BcI</u>I site at the stop codon of the Fd chain provided a convenient method for removal of all non-yeast 3' untranslated DNA. This was accomplished using the strategy outlined in Figure 34, and generated the plasmid, DNG 1428.

[0254] Since the stop codon was introduced into the hinge region by site-specific mutagenesis of a heavy chain fragment cloned into M13, the possibility existed that unwanted mutations could have been introduced during the mutagenesis step. To ensure that such mutations were not present, an Fd gene fused to the invertase signal sequence and shortened PGK promoter and consisting of known coding sequences was constructed using the strategy outlined in Figure 34, generating the plasmid, pING 14444.

(4) Construction of yeast expression plasmids containing the chimeric L6 Fd gene from plNG1444 fused to the PGK polyadenylation signal.

40 [0255] In order for yeast to produce an intact, functional Fab molecule, a balanced synthesis of both light and Fd-chain proteins must occur simultaneously within the cell. As described in Example V, one approach is to place the light chain and Fd genes on separate shuttle vectors containing separate selective markers and to transform these vectors into a yeast strain detective for both selective markers.

[0256] The Fd gene from pINS11444 (Figure 34) was cloned as a BamHI-Yoot fragment into two medium copy number yeast= f_coei institute vectors containing sequences for registeation in yeast and the PGK polyadenylation, transcription termination signal: pINS604CVS for leg2 selection and pINS1150 for leg3 selection (see Figures 29, 30). The two plasmides resulting from these constructions—JNR1445 (urs) and pINS1446 (suc2) are shown in Figure 3 and pINS1450 for leg3 and pINS1446 (suc2) are shown in Figure 3 and pINS1450 for leg3 and pINS1450 for

(5) Secretion of chimeric L6 Fab from transformed yeast cells.

50

[0257] Two separate transformation experiments were performed in an attempt to obtain both light and Fd-chain synthesis in yeast cells. Four µg each of plNG1445 (Figure 35) and plNG1441 (Figure 30) and separately of plNG1445 (Figure 36) and plNG1442 (Figure 37) and plNG1442 (Figure 38) were co-transformed into <u>S. cerevisiae</u> strain BB331c (MATa, <u>ura3</u>, <u>ura2</u>) by selecting for growth on SD agar (2% plucose, 0.67% yeast nitrogen base, 2% agar). Ura* Leu* transformants appeared at two to three days of incubation at 30°C.

[0288] Five colonies were inoculated from each plate into 6 ml SD broth supplemented with 50 mlM sodium succinate, pH 5.5, and grown for 65 hours at 30°C. The cells were removed by centrifugation and analyzed by ELISA for the levels of light chain. The results of these assays revealed that the levels of light chain in the culture supernatants of

the pING1448 + pING1414 transformants were three to six times higher than the levels in the culture supernatants of the pING14445 + pING1414 transformants. The culture supernaturate for each propup of transformants were next concentrated by ultrafiltration on a Centricon 30 filter (Amicon Corp.) and run on a 10% polyacrylamida gel under non-educing conditions. The proteins were bitted to nitrocellulose paper and probed with goat anti-thuran kappa ariserum followed by peroxidase-labeled rabbil-anti-poat antiserum. The concentrated supernatant from the pING1445 and piNG1445 transformants contained a significant anti-kappa cross-reactive ener over a large portion of the bld with only a faint cross-reactive band at the position expected for the Fab protein. By comparison, the concentrated supernatant from pING1445 pING1441 transformants contained retailively filtils ensered anti-thurant kappa cross-reactive protein on the blot. In addition, one of the five samples (No. 4) contained an especially intense, distinct anti-kappa cross-reactive band with which will revise the protein of the blot. In addition, one of the five samples (No. 4) contained an especially intense, distinct anti-kappa cross-reactive band with migrated at the costilion excented for an Fab protein.

(6) Purification of chimeric L6 Fab from yeast culture supernatant.

[0259] To establish that the Fab-size anti-kappa cross-reactive protein secreted by the yeast is indeed L6 chimeric Fab protein required the purification of sufficient quantities for performance of binding assays. The pING1441 + pING1445 transformant isolate No. 4 was, therefore, grown in one liter of SD broth supplemented with 50 mM sodium succinate, pH 5.5, for 95 hours at 30°C. The cells were removed by centrifugation and the culture supernatant was analyzed by ELISA for the level of light chain protein. The supernatant contained approximately 130 µg/L of light chain protein. The culture supernatant was next concentrated by ultrafiltration over an Amicon YM30 filter to 20 ml. The concentrated supernatant was washed with 130 ml 10 mM potassium phosphate, pH 7.5 (buffer A) and re-concentrated over the YM30 filter to 12.5 ml. The concentrated supernatant was next brought to 54 ml with buffer A and loaded onto a 1.5 ml S-Sepharose column equilibrated with buffer A. The column was washed with 20 ml buffer A and subjected to a linear gradient of 0 to 200 mM sodium chloride in buffer A (40 ml total volume). ELISA analysis of the column fractions revealed a large anti-kappa cross-reactive peak between fractions 8 and 21 corresponding to a salt concentration of approximately 60 mM. These fractions were pooled, concentrated on Amicon YM10 and Centricon-10 filters (Amicon Corp.) to 51 µl and analyzed on non-reducing and reducing polyacrylamide gels using coomassie blue staining and Western blotting with anti-human kappa and anti-human Fab antisera. These analyses revealed an essentially pure protein which migrated at approximately 46 kd on the non-reducing gel and resolved into two bands running at approximately 23 and 24.5 kd on the reducing gel which corresponds to the predicted (based on amino acid sequence) molecular weights for light chain and Fd proteins, respectively. The smaller of the two bands strongly reacted with antihuman kappa antiserum on the Western blot. Both of the protein bands reacted with anti-human Fab antiserum on the Western blot.

(7) Studies performed on the chimeric L6 Fab secreted by yeast.

[0260] The primary activity of an Fab molecule is its ability to bind to the target antigen. The yeast-derived chimeric Fab was, therefore, tested for its ability to bind directly to an L6 antigen-positive cell line and for its ability to inhibit binding of mouse L6 antibody to antigen-positive cells.

[0251] <u>Direct Binding Assay.</u> Cells from the human colon carcinoma cell line 3347, which contains the L6 antigen at the cell surface, were used as a largets. Cells from the antigen-negative cell line, T51, were used as a negative control. The target cells were first incubated for 30 minutes at 4°C with either yeast-derived chimeric L6 Fab, 592/C cell-derived chimeric L6 antibody, or with mouse L6 antibody. This was followed by incubation with F1TC-dabelled goat anti-human kappa immunoglobulin for the chimeric Fab, F1TC-dabelled goat anti-human lgG for chimeric antibody, or with F1TC-babelled goat anti-mouse immunoglobulin for the mouse antibody. Both labelled antibodies were obtained from TAGO (Burlingame, CA) and used at a dilution of 1:50. Antibody binding to the cell surface was determined using a Coulter Model EPIC-C cell storfer.

[0262] As shown in Table 12, the yeast-derived chimeric L6 Fab bound to the L6 positive 3347 line. The yeast-derived chimeric L6 Fab did not bind above background to the L6 negative T51 line.

[0263] Inhibition of Binding. As the next step, we studied the extent to which graded doses of the yeast-derived of inhibit binding of an FITC-labelled mouse L6 antibody to the surface of antibon positive colon carcinoma 3347 cells.

[0264] The yeast-derived chimeric L6 Fab inhibited the binding of the directly labeled mouse L6 antibody. A higher concentration of the yeast L6 Fab, however, was required to achieve 50% inhibition of mouse L6 antibody binding the target-ells than was required for the same degree of binding inhibition by Sp2/O cell-derived chimeric L6 antibody.

(8) Conclusions

35

[0265] A process is disclosed by which yeast can be genetically engineered to secrete functional Fab domains of

immunoglobulins. The yeast-derived chimeric Fab in this example binds to the appropriate target antigen. Such Fab molecules provide convenient targeting agents for a variety of diagnostic and therapeutic uses. This process also demonstrates the feasibility of secretion of heteroloous heterodimeric molecules from vessor.

TABLE 12

		IABLE 12				
	BINDING ASSAYS OF CHIMERIC LE	FAB PRODUCED BY YEAST ON AN ANTIGEN-NEGATIVE CELL LINE	L6 ANTIGEN-POSITIVE AND AN L6			
	Binding Ratio ^b for:					
- [Antibody ^a	3347 Cells (L6 +)	T51 Cells (L6-)			
Γ	Sp2/O Chimeric L6	103	1			
	Yeast Chimeric L6 Fab	32	1			

a. All antibodies were used at a concentration of 10 µg/ml.

10

15

20

b. The binding ratio is the number of times brighter a lest sample is than a control sample treated with FITC-conjugated second antibody. Goal anti-human antibody was used as the second antibody for the Sp2/O chimeric L6 antibody and goal-anti-human kappa antibody was used as the second antibody for the vesst Fab.

EXAMPLE VII: Secretion of Functional Chimeric Fab Molecules From Bacteria

[0266] Bacteria are suited to production of chimeric antibodies expressed from mammalian cDNA since entire coding sequences can be expressed from well characterized promoters. <u>Escherichia col</u> is one of many useful bacterial species for production of foreign proteins (robland, B., gl <u>al.</u>, <u>BloTachnology</u> 4427 (1995)), since a wealth of genetic information is available for optimization of its gene expression. <u>E. col</u> can be used for production of foreign proteins internally or for ascretion of proteins out of the cytoplasm, where they most lotten accumulate in the periplasmic space (Gray et al., Gene <u>98</u> 247 (1985); Oka <u>et al.</u>, <u>Proc. Natl. Acad. Sci. USA <u>82</u> 7212 (1985)). Secretion from the <u>E. col</u> cytoplasm has been observed for many proteins and requires a signal sequence. Proteins produced internally in bacteria are often not folded properly and precipitate into subcellular particles called inclusion bodies (Schoner <u>et al.</u>, <u>BioTechnology</u> 3:15 (1985)). Protein secreted from bacteria, however, is often folded properly and assumes native secondary and-terilary structures (Hstung <u>et al.</u>, <u>BioTechnology</u> 4:991 (1986)). Although immunoglobulin peptides have been synthesized in genetically engineerad <u>E. col</u> (Cabilly <u>et al.</u>, <u>Proc. Natl. Acad. Sci. USA</u> 31:5399 (1984), Llu <u>et al.</u>, <u>Proc. Natl. Acad. Sci. USA</u> 31:3399 (1984), Llu <u>et al.</u>, <u>Proc. Natl. Acad. Sci. USA</u> 31:5399 (1984), Llu <u>et al.</u>, <u>Proc. Natl. Acad. Sci. USA</u> 31:5399 (1984), Llu <u>et al.</u>, <u>Proc. Natl. Acad. Sci. USA</u> 61:5399 (1984), Llu <u>et al.</u>, <u>Proc. Natl. Acad. Sci. USA</u> 61:5399 (1984), Exes <u>et al.</u>, <u>Nucl. Acid. Res.</u> 12:2791 (1984)), there are no reports of secretion of these peptides from E. col is at functional antibocies or antibody fragments.</u>

[0287] An Fab molecule consists of two nonidentical protein chains inked by a single disulfide bridge. These two chains are the intact antibody high chain and the V, J, and C₁₁, portions of the antibody heavy chain, Fd. The procept cDNA clones for the L6 chimeric light and Fd gene have already been identified. In this example, these cDNA clones were organized into a single bacterial operon (a distitution inseesage) as gene fucions to the pactital tysas (pall) gene leader sequence from <u>Erwinia carotoxora (Lei et al.</u>, J Bacteriol., in press (1987)) and expressed from either of two strong, regulated promotes. The result lis a system for the simultaneous expression of two protein chains in <u>E</u>. Coll, and the secretion of immunologically active, properly assembled Fab of L6 chimeric antibody into the culture growth notes.

A. Construction of E. coli expression systems for L6 Chimeric Fab.

1. Assembly of the pelB leader sequence cassette.

[0268] Envinia carotovora EC codes for several pectate lyases (polygalacturonic acid trans-eliminase) (Lei et al., Gene 35.63 (1985)). Three pectate lyase genes have been cloned, and the DNA sequence of these genes has been determined. When cloned into E. coli under the control of a strong promoter, the pell gene is expressed and large quantities of pectate lyase accumulate in the periplasmic space. The pell signal sequence functions efficiently in E. coli and was used as a secretion signal for antibody genes in this example. The nucleotide sequence surrounding the signal sequence of the pell gene is shown in Figure 36a.

[0259] The pall signal sequence contains a Heall restriction site at amino acid 22, adjacent to the signal peptidase cleavage site a lae-lae Plasmid pSS 1004 (Lie if et al., <u>J. Bacteriot</u>, in press (1987), containing the pell gene in plasmid vactor pUC8 (Vieirra and Messin, <u>Gene 19:258</u> (1982)), was digested with <u>Heall</u> and <u>EcpRI</u>. This DNA was ligated with an eight base pair <u>Sell</u> infect to <u>Sell</u> and <u>EccPI</u> cut pBR322. The resulting plasmid contained a 300 by fragment which included the 22 amino acid leader sequence of pelB and about 230 by of upstream <u>E. caratovora</u> DNA. This pismid pING173, contains an insert that upon dispession with Stat and treatment with T4 DNA polymerase can

ligated directly to a DNA fragment flanked by the first amino acid of a mature coding sequence for any gene to generate a protein fusion containing a functional bacterial leader sequence in frame with the incoming gene. The <u>Staf</u> to <u>Ecoplin</u> restriction fragment in piNG173 was cloned intop UC18 (Yanich-Perro<u>n et al., Gene S3</u> 103 (1985)) to generate pFR175, which contains the <u>pelB</u> leader and adjacent upstream non-coding sequence (including a ribosome binding site) downstream of the lac promoter. The construction of pR1715 is outlined in Figure 396.

2. Preparation of chimeric L6 light gene for bacterial expression.

[0270] The intact L6 chimeric light chain gene containing an Agtil restriction site at the signal sequence processing all and a unique Bgill site downstream of the gene was excised from the yeast expression plasmid plNG 1298 (Figure 25a) as a 1200 bp DNA tragment. This fragment was inserted into plasmid pFR175. The resulting plasmid, pRR177-8, contained an in-frame fusion of the pellg leader and the L6 light chain downstream of the lag promoter residing in the parent plasmid. A number of derivatives of this plasmid were constructed to delete noncoding sequences from both the 5' and 3' endes to the pellg-light chain gene tusion in pFR177-8. Upstream noncoding sequences were deleted making use of an Neel restriction site at 1-48 bp from the pell seader sequence initiation codon (Figure 35) generating pRR180-2. The 3' noncoding sequences were eliminated by substituting a fragment from the plasmid optimized for L6 light chain expression in yeast, pINSI 1431 (see Figure 27a), into pRR179 is generate pRR191. Another plasmid, pRR180, is similar to pFR191 but contains 90 bp of noncoding eukaryotic DNA at the 3' end of the light chain gene. Those constructions are shown in Figure 37.

3. Preparation of chimeric L6 Fd gene for bacterial expression.

20

35

50

[0271] The intact L6 chimeric Fd gene containing an <u>Sst</u>1 restriction site at the signal sequence processing site, a <u>Sct</u>1 site introduced by site directed mutagenesis (Figure 32a, b) and creating a termination codor at amino acid 226, and a unique <u>Bam</u>HI restriction site downstream of the gene was excised from the plasmid pING1406 (Figure 32) as a 860 bp DNA fragment. This DNA fragment was inserted into plasmid pFRI175 generating an in-frame fusion of the <u>pBIS</u> leader sequence and the L6 Fd gene downstream of the <u>pice</u> promoter, <u>PRI178-5</u>. A number of derivatives were constructed to delete noncoding sequences from both the 5' and 3' ends of the sequence contained in <u>PRI178-5</u>. The 3' noncoding sequences were eliminated by substituting a restriction fragment from the plasmid optimized for L6 Fd expression in yeast, pING1426 (Figure 34), which contains an <u>xhol</u> linker immediately following the termination codor of the Fd gene, generating plasmid pFRI186. Removal of <u>E. caratovora</u> DNA sequences upstream of the <u>Nidel</u> site at 48 from the leader sequence generated plasmid pFRI186. The construction of these plasmids is shown in Figure 33.

4. Multicistronic expression system for light chain and Fd gene.

[9273] For production of bacterially derived Fab, both light chain and Fd need to be produced simultaneously within the cell. Using the plasmide constructed with each of these genes separately, a series of asyression vectors were constructed that contain both genes aligned so that transcription from a single promoter will specify both genes. This was done in a way that minimized the noncoding DNA between the two genes to 60 bp. Each gene has a riboome binding alte needed for translation initiation and the identical DNA sequence from -48 to the gells leader: antibody genes in plantices. Several cloning steps were required to align the two genes together. A portion of the light chain gene light data (see the gells leader in pRFI180-2 was cloned downstream of the Fd gene in pRFI186 to generate pFK100. The remainder of the light chain gene was subcloned into pFK100 from pRFI190 and pRFI191 were cloned into pFK101 generatins containing 3° deletions of eukaryotic sequences from pRFI190 and pFRI191 were cloned into pFK101 generating pFK102 respectively. DNAI regressions from pFRI192 and pFK101 were ligated to generate pFK101 stripe from the Fd gene. Maps of the Fd and light chain gene cassettes in these plasmids are shown in Flaure 93.

5. Placement of the dicistronic message for light chain and Fd under the control of inducible promoters.

[0273]. Plasmids pFK101, pFK102, pFK103, and pFK104 contain Fd and light chain genes cloned sequentially under the control of the lag promoter in vector pUC18 or pUC19. In <u>E. coll</u> strains such as JM103 FlagaC (Messing <u>et al.</u> Nucl. Acids. Fles. <u>9</u>:309 (1981)), the amount of light chain that accumulates in the periplasm is not affected by the <u>lice</u> promoter inducing agent isopropi B-D-thiogalactopyranoside (IPTG), see Table 13. In addition, bacterial growth is slow-er (compared to cells containing pUC19), and bacterial colonies exhibit an aftered morphology being small, dynd rough, suggesting that constitutive foreign gene expression is deleterious to cell growth. Two strategies were used to piece this cone cassetie under more light in requisited promoters.

[0274] First, a Psti to EcoRi fragment from pFK104 was ligated to pIT206 to place the Fd and light chain gene cassette

under the direct control of the <u>Salmonella typhimurium araB</u> promoter, a well characterized, strong promoter in <u>E. coli.</u>
A restriction map of pT206 and construction of pT104 is shown in Figure 40. Use of the <u>araB</u> promoter and its requirage
protein <u>ara</u>C for the expression of bacterial genes is described in U.S. Patent Applications 695,309 filled January 28,
1995, and 797.472, filed November 13, 1995. As is seen in Table 14, the resulting plasmid, pT1104, is now regulated
for the synthesis of light chain by the addition of arabinose to the culture growth media. At least 10 floid induction is
effected by arabinose addition. Although Fab secreted into the growth medium increases more than 10 flot, cell growth
stops after induction with arabinose. This confirms that high level expression of the Fab genes is deleterious cell
growth. Becterial colonies hathoring pT104 are phenotypically indistinguishable from <u>E. coli</u> harboring pT1206 when
grown in the absence of arabinoses.

[0275] Second, a DNA fragment containing the <u>laci</u> gene, a repressor of the <u>lac</u> promoter, was cloned into the high copy expression vector pFK102. Expression of <u>laci</u> from a high copy number vector is useful to requisite expression of the <u>lac</u> promoter on a high copy number vector (Flussel <u>et al.</u>, Pleasmid, press (1987). Having et al., <u>Biotechnology</u> 4:991 (1986)). A 1.7 kb <u>EcoPI</u> tragment containing the <u>laci</u> gene on pMC9 (Calos <u>et al., Proc. Natl. Aced. Sci. USA 80</u>; 0315 (1989)) was excised, flused in with 14 polymerase to butten desl, <u>lagrad with Pst</u> linkers and cloned into the unique of the <u>laci</u> sist of pFK102 to generate pFK102<u>laci</u>. The map of pFK102<u>laci</u> is shown in Figure 40b. The selection procedure used to identify the correct clone assured that the resulting plasmid, pFK102<u>laci</u>. contained a functionally repressed <u>lac</u> promoter. All white or light pink colonies on MacConkey-lactose pitates contained plasmids with <u>laci</u> inserts white transformants containing pFK102_L alone were red, indicating the functional repression of the lac promoter by the high copy number laci gene. Table 14 shows that expression to bacterial Fab trone elle containing pFK102_L is similar to expression from pFK102. Unlike cells containing pFK102_L is similar to expression from pFK102_L thick collections entaining pbK102_L is containing pFK102_L is similar to expression from pFK102_L containing pbCK102_L is containing pCK102_L is contai

B. Expression, SDS-PAGE, and Purification of Bacterially Produced Fab

Growth of E. coli harboring cloned antibody genes.

[0276] Plasmid DNA was transformed into either <u>E. coli</u> JM103 or MC1061 by standard <u>E. coli</u> transformation procedures. Bacterial cultures were grown in TYE (tryptone 1.5%, yeast extract 1.0%, and NaCl 0.5%) supplemented with the appropriate antibiotics (pencillalle 250 ug/ml, letteracyline 15 ug/ml). Bacterial cultures were grown in volumes of 5 ml to 1 liter at 37°C to an optical density OD600 = 0.8 (approximately 4.X 108 cell/ml) and aliquots were induced with IPTG (0.2 mM), lactose (1.0%), or arabinose (1.0%). Cultures were grown for an additional time period of 4 to 21 hr. Portions of each culture were analyzed for light chain production. Protein was released from the periplasmic space of <u>E. coli</u> cells by osmotic shock as described (Yanagots et al., J. <u>Bacteriol</u> 168:937 (1986)). Alternatively, culture supermatents were assayed directly for the presence of antibody chains.

(2777) Quantization of L6 light chain was by ELISA with goat anti-human Kappa light chain antibody (Cappal, Malwam, PA). F1 could be detected by ELISA with mouse monoclonal anti-human F4 antibody (Cabibichem, San Diego, CA). Table 13 shows representative data for expression of light chain reactive material in <u>E_coll</u> periplasmic extracts. Light chain is secreted from the bacterial cytoplasm into the purplam. Antibody chains a real also released from the bacterial into the culture supernstant. This is an unusual discovery and may be a unique property of the L6 Fab among avaraycic profelse expressed in <u>E_coll</u>. Under certain contions, however, bacterial proteins are known to be released from <u>E_coll</u>. Old Abrahmsen <u>et al.</u>, Nucl. Acids Res. <u>14.7487</u> (1986), Reges et <u>al.</u>, <u>JB action [1986</u>), Table 14 compares the amount of light chain secreted into the periplasm with the amount secreted into the culture supernstant. Light chain reactive material is present in plasmid containing cultures harboring cloned light chain alone or light chain plus Fd. The bet producers of Fab (pFK102, pFK104, and pFK102<u>led</u>) typically secrete 300 - 1000 ng/ml of ELISA reactive light chain into the culture model. A separate construct was made in which the light chain get is followed by the Fd gene (pFK107). This construct directs synthesis and secretion of Fab at similar levels to the constructs with the genes in the inverse order. Thus, the erean order is not critical for secretion of Fab at similar levels to the constructs with the genes in the inverse order. Thus, the erean order is not critical for secretion of Fab.

2. SDS-PAGE of bacterially produced chimeric L6 light chain and Fd.

50

[0278] Bacterially produced antibody chains were analyzed by polyacrylamide gel electrophoresis under reducing and non-reducing conditions. Protein extracts of tysed whole bacterial cells, protein released from the periplasmic space by osmotic shock, and protein secreted into the culture supernatian were analyzed electrophoretically. Transfer of gel separated protein under full reducing conditions to nitrocellulose and immunological staining with goat anti-human light chain antibody by Western analysis revealed that a protein of the same molecular weight as authentic L6 chimaric light chain was present (about 23 Kd). Analysis of protein samples by SDS-PAGE under non-reducing conditions showed that extracts from cells harboring a plasmid with the light chain gene alone (pHR191 or pRR190) crained a large proportion of the light chain rescrive material associated into a higher molecular weight form. Much of

this material ran at about 46 Kd in what is likely to be a light chain dimet. Light chain dimers have been observed from myelcane cells producing only light chain. There are also other immunoreactive protein bands that may represent non-specific disufficient formation between light chain and <u>E. coil</u> proteins. Protein samples (periplasmic extracts or culture supermatants) from <u>E. coil</u> cells harboring both the light chain and the Figenes contain a light chain receive band at about 48 Kd when separated under non-reducing gel conditions which runs at a slightly higher noticular weight than the bacterial light chain dimer. This material is bacterially produced L6 chimeric Fab. In <u>E. coil</u> harboring PFK 102 lact, PFK101, PFK102, PFK103, or PFK104 the 48 Kd band observed on an SDS gel run under non-reducing conditions is the most prominent immunoreactive specie. In addition, the background smear of immunoreactive proteins seen in extracts containing the light chain only is greatly reduced in extracts from cells containing be light chain and Fig.

3. Purification of bacterially produced chimeric L6 Fab.

[0279] Immunologically and functionally active (see below) bacterial Fab was purified from either culture supernatants or periplasmic protein extracts of <u>E. coli</u> harboring pFK102<u>laci</u> or pIT104. For purification of periplasmic material, the periplasmic fraction from 1 liter of cells induced for 4 hours was released into 50 ml of distilled water. This material was centrifuged for 20 minutes at 5000 g and filtered through a 0.45 jum filter. The periplasmic extract was then concentrated over a YM10 membrane (Amicon) to about its ml. This material was dituted 8 fold into starting buffer (10 mM RZHPO4, pH 7.5) and applied to a 1 ml S-Sepharose column at a flow rate of 1.0 ml/min. The column was washed with 25 ml of starting buffer and eluted with a 0 to 200 mM NaCl gradent in starting buffer (200 ml total volume). The immunoreactive gradent peak was pooled (elution was at about 100mM) and concentrated on a Centricon 10. Purified Fab was stored

in PBS + 2.0% BSA.

[288] For purification of secreted Fab from 1 liter of bacterial culture supermatant, the cells were removed by centrifugation after growth for 21 hours with inducing agents and the supermatant was filtered through a 0.45 µm filter. The media was concentrated over a W101 membrane (Amicon) to about 15 mt, then diluted with 10 mM (21PD-04 to 105 mt. This material was applied to a 1.6 mt. S-Sepharose column and eluted with a 0 to 200mM NaCl gradent in 40 mt. Pab recovered from S-Sepharose chromatography was greater than 70% pure as determined by densitionerly to 4 mt. Pab recovered from S-Sepharose column and eluted with a 0 to 200mM NaCl gradent in 40 mt. Pab recovered from S-Sepharose column and eluted with a 0 to 200mM NaCl gradent in 40 mt. Pab recovered from S-Sepharose column and eluted with a 0 to 200mM NaCl gradent in 40 mt. Pab recovered from S-Sepharose column and eluted with a 20 mt. Pab recovered from the central culture supermatants resolves into two major protein bands of about 23 Kd and 42.5 Kd on at 15% reducing get. The melecular weight of 2 kd and 42.5 Kd on a 15% reducing get. The melecular weight of 2 kd and 42.5 Kd on a 15% reducing get. The melecular weight of 2 kd and 42.5 Kd on a 15% reducing get. The melecular weight of 2 kd and 42.5 Kd on a 15% reducing get. The melecular weight of 2 kd and 42.5 Kd on a 15% reducing get. The missing of 3 kd and 42.5 Kd on a 15% reducing get. The missing of 3 kd and 42.5 Kd on a 15% reducing get. The substitute supermatants are indictinguishable by all analytical Criteria is seted free.

4. Functional binding activity of bacterially produced chimeric L6 Fab to the L6 antigen.

[0281] Bacterially produced Fab purified by S-Sepharose chromatography was tested for binding to Le antigen containing cells. As ehown in Table 145, bacterial Fab binds specifically to the human colon acrinoma cell line 3347. Cells from the T cell line T51 were used as a negative control. Target cells were incubated for 30 minutes at 4°C with bacterially produced Le chimeric Fab, intact L6 chimeric ranking to produced in \$920 cells, or mouse L6 antibody purified from ones ascites. This was followed by incubation with FITC-labeled goat anti-human light chain antibody for Fab detection, FITC-labeled goat anti-human immunoglobulin for mouse antibody detection, or with FITC-labeled goat anti-mumirine immunoglobulin for mouse antibody detection, or with FITC-labeled goat anti-mumirine immunoglobulin for mouse antibody detection, or with FITC-labeled goat anti-mumirine immunoglobulin for mouse antibody detection, or with FITC-labeled goat anti-mumirine immunoglobulin for mouse antibody detection, or with FITC-labeled goat anti-mumirine immunoglobulin for mouse antibody detection, or with FITC-labeled goat anti-mumirine immunoglobulin for mouse antibody detection, or with FITC-labeled goat anti-mumirine immunoglobulin for mouse antibody detection, or with FITC-labeled goat anti-mumirine immunoglobulin for mouse antibody detection, or with FITC-labeled goat anti-mumirine immunoglobulin for mouse antibody detection, or with FITC-labeled goat anti-mumirine immunoglobulin for mouse antibody detection, or with FITC-labeled goat anti-mumirine immunoglobulin for mouse antibody detection, or with FITC-labeled goat anti-mumirine immunoglobulin for mouse antibody detection, or with FITC-labeled goat anti-mumirine immunoglobulin for mouse antibody with FITC-labeled goat anti-mumirine immunoglobulin for mouse antibody with FITC-labeled goat anti-mumirine immunoglobulin for mouse antibody with FITC-labeled goat antibody with FITC-labeled goat antibody with FITC-labeled goat antibody with FITC-labeled goat antibody with FITC-labele

[2282] Bactorially produced Fab also exhibits characteristic binding inhibition of FITC-labelled mouse L6 antibody to the surface of antipen positive 3474 colon carcinome cells. Beaterially produced Fab and Sp2/D derived chimeric L6 have similar binding inhibition profiles, thereby suggesting that the avidity of bacterially produced Fab and Sp2/D derived chimeric L6 are similar.

Conclusions 50

10

35

[0283] A process is disclosed whereby <u>E. coli</u> has been used as a host to produce functionally active Fab domains of immunoglobulins and to secrete these into the periplasmic space and also in the culture medium. This molecule exhibits binding properties expected of a property assembled antibody recognition site. This technology can be used to express antibody genes with other binding specificities in E. coli.

- 1. Proteins encoded by modified antibody cDNA clones can be secreted from bacteria using a signal sequence
- 2. Two antibody genes can be expressed from a single bacterial promoter as a dicistronic message.
- 3. Two foreign proteins (in this example antibody light chain and Fd) can assemble properly, i.e., assume correct

secondary, tertiary, and quaternary structure when secreted from bacteria.

- 4. At least two, and probably many bacterial promoters can be used for expression of antibody genes.
- 5. This example is a general method whereby genes encoding other antibody chains can be expressed together as a dicistronic message; these include either light chain and Fd genes or light chain and intact heavy chain genes.
- 6. The gene order with respect to the promoter is not important in the ability of <u>E. coli</u> to produce Fab. A construct of the Fd gene followed by the light chain works as well as the genes organized in the inverse order.
- 7. Fab can be secreted from <u>E. coli</u> into the culture supernatant where it is stable and can be purified. Most Fab chains that pass the cytoplasmic membrane are secreted into the culture supernatant.

Microorganism Deposits

5

15

20

25

30

35

40

45

50

55

[0284] Saccharomyos cerevisiae BB331C (41/42-5), G187 was deposited at the ATCC on July 9, 1997 and given access number 20856. Escharichia coil JM 103 (pFK1021 aci), G188 was also deposited therein on the same date and given access number 67457. Both deposits were under the Budapest Treaty.

TARIF 13

ſ	7	П	٦	_	_		_	Т	
		ng/ml of culture	+	28	88	ð.	88		
	LI PERIPLASM	o lm/gu		38	89	88	16		
MOLE 13	QUANTITATION OF LIGHT CHAIN FROM E. COLI PERIPLASM	plasmid		pFK101	pFK102	pFK103	pFK104		
	QUANTITATION OF LI	ng/ml of culture	*	0	=	412	241	77.5	
		ē			8.5	399	200	463	
		plasmid		oRR175	0HR177-8	DBR180	pHR190	pRR191	

E_coil_JANTQ or MC1061 (results similar) was transformed with each plasmid. Fresh transformants were cultured in TYE at 37 °C to an ODR00 = 0.B. Cultures were included and activate (FVE) was not grown at 37 °C to 4 future. Periplasmid protein entertas were prepared, included and an adjustance were an an adjustance were instanced were transfer or EUSA with those tan inmant facpor antitority. Each value is the servage of at least two separates experiments. Removal or non-coding sequences both 6" and 3" to the amboory gave effected in increase on light chain accumulation in the periplasm.

TABLE 14

plasmid	Inducer	edif.	Supemalant	icad	Dacipleom
		2	The state of the s	181	III
		4 hr	21 hr	+	21 hr
pRR180		0	þ	90	5
pRR190		'n	28	241	2 2
pFK102		12	2		? 'E
pFK102	•	25	828		! \$
pFK104		52	2	: 6	? ?
pFK104	•	150	230	. 99	, K
pFK102laci		25	380	: %	3 5
pFK102laci	•	72	909	37	3 4
plT104		13	2	, ç	; }
pIT104	•	150	216	. 6	2 15
Plasmid containing E. coli s	trains were grown, prepared	d, and assayed as described	in Table 13. For pRR190, pf	Plesmid containing E. coli strains were grown, prepared, and assayed as described in Table 13. For pRR190, pFK102, pFK102, and pFK102laci cells were induced with	aci cells were induced with
bacteria were removed by	nduced with 1% arabinose.	Each value is the average of	at least two separate experi	0.2 mm Int. (c) prilve was induced with 'N additions. Each will use the tevelop of at least two separate experiments. For analysis of E. coal custure supernatants, hadrests were named to remain any contributional to remain relative supernatants.	culture supernatants,
	no company or management	and passed and supplied	ugil a 0.45 um IIIBI. Yalues	are expressed in ng/mi of cul	Ture.
nd - not determined					

TABLE 15

	Binding ratio*				
Antibody	3347 cells L6 +	T51 cells L6-			
Standard mouse L6	95	1			
Sp2/0 chimeric L6	116	1 1			
Bacterial L6 Fab	54	1 1			
Standard L6 Fab	16	1 1			

^{*} The binding ratio is the number of times brighter a test sample is than a control sample treated with FITC-conjugated second antibody.

Claims

10

15

20

an

45

- 1. A polynucleotide molecule comprising a prokaryotic promoter region in operable linkage to a dicistronic transcription unit, the unit encoding a heavy chain immunoglobulin, or a fragment thereof, and a light chain immunoglobulin, the heavy and light chains being separately operably linked to a sequence coding for a polypeptide secretion signal.
 - 2. A molecule according to Claim 1 wherein the heavy and light chains are chimeric.
- 3. A vector comprising recombinant DNA and a polynucleotide molecule as defined in any of Claim 1 or Claim 2 and capable of expressing the polynucleotide molecule in a prokaryotic host.
 - 4. A prokaryotic host capable of producing an immunoglobulin transfected with a vector as defined in Claim 3.

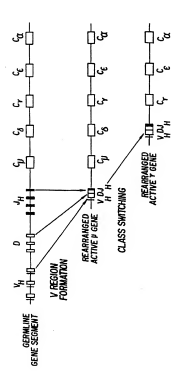
Patentansprüche

- 1. Polynucleotidmolekül, welches eine prokaryotische Promotorregion in funktioneller Verknüpfung mit einer dicistronischen Transkriptionseinheit aufweist, wobei die Einheit eine schwere Kette eines Immunglobulins oder ein Fragment davon und eine leichte Kette eines Immunglobulins codiert, wobei die schweren und leichten Ketten getrennt funktionell mit einer Sequenz verknüpft sind, die Polypeptidsekretionssignal codiert.
 - 2. Molekül nach Anspruch 1, bei welchem die schweren und die leichten Ketten chimär sind.
- 3. Vektor, welcher rekombinante DNA und ein Polynucleotidmolekül gemäß der Definition in einem der Ansprüche 1 oder 2 enthält, und welcher fähig ist, das Polynucleotidmolekül in einem prokaryotischen Wirt zur Expression zu bringen.
 - 4. Prokaryotischer Wirt, welcher in der Lage ist, ein Immunglobulin zu erzeugen, welcher mit einem Vektor gemäß der Definition in Anspruch 3 transfiziert ist.

Revendications

- 1. Molécule polynucléotidique comprenant une région promoteur provenant d'un procaryote, en liaison opérationnelle avec une unité de transcription dicistronique. l'unité codant pour une immunoglobuline à chaîne lourde ou un fragment de celle-ci, et une immunoglobuline à chaîne légère, les chaînes lourde et légère étant liées séparément de manière opérationnelle à une séquence codant pour un signal de sécrétion de polypeptide,
- 2. Molécule suivant la revendication 1, dans laquelle les chaînes lourde et légère sont chimériques. 55
 - 3. Vecteur comprenant un ADN recombinant et une molécule polynucléotidique comme défini dans l'une quelconque des revendications 1 et 2, et capable d'exprimer la molécule polynucléotidique dans un hôte procaryote.

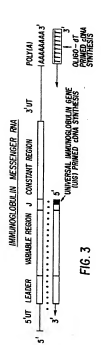
	4.	Hôte procaryote capable de produire une immunoglobuline, transfecté par un vecteur défini dans la revendication 3.
5		
3		
10		
		·
15		
20		
25		
		•
30		
35		
35		
40		
45		
50		
55		

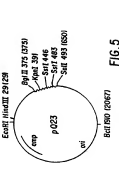


- 2

- 8	leavy chain 3-C region	
hum	in heavy chain J regions	CHI
JH 1	GCTGAATACTTCCAGCACTGGGGCCAGGGCACCCTGGTCACCGTCTCCTCAG	
JH2	CTACTGGTACTTCGATCTCTGGGGCCGTGGCACCCTGGTCACTGTCTCCTCAC	
JH3	ATGCTTTTGATGTCTGGGGCCAAGGGACAATGGTCACCGTCTCTTCAG	
JH4	ACTACTTTGACTACTGGGGCCAAGGAACCCTTGGTCACCGTCTCCTCAG	
JHS	ACACTGGTTCGACTCCTGGGGCCAAGGAACCCTGGTCACCGTCTCCTCAG	
	AT(TAC) COTATION COTOTOGOGOGO ANGGARCOU TOGTO ACCOTOTOCTORO	
Car	AT(TAC) 5GGTATGGACGTCTGGGGGCAAGGGACCACGGTCACCGTCTCCTCAG	
Con	rensus TCGACCTCTGGGGCCAAGGAACCCTGGTCACCGTCTCCTCAG	
mou:	se heavy chain J regions	
		CHI
JH 1	TACTGGTACTTCGATGTCTGGGGCGCAGGGACCACGGTCACCGTCTCCTCAG	
JH2	TACTTTGACTACTGGGGCCAAGGCACCACTCTCACAGTCTCCTCAG	
JH3	CCTGGTTTGCTTACTGGGGCCAAGGGACTCTGGTCACTGTCTCTGCAG	
JH4	TACTATGCTATGGACTACTGGGGTCAAGGAACCTCAGTCACCGTCTCCTCAG	
Con	TTTGACTACTGGGGCCAAGGCACCACGGTCACCGTCTCCTCAG	
Ig	ight chain J-C region	
hum	n Kappa J region	
	J1C	
	GGACGTTCGGCCAAGGGACCAAGGTGGAAATCAAAC	
JK2	ACACTTTTGGCCAGGGGACCAAGCTGGAGATCAAAC	
JK3	TCACTTTCGGCCCTGGGACCAAAGTGGATATCAAAC	
	TCACTTTCGGCGGAGGGACCAAGGTGGAGATCAAAC	
JK5		
Con:	TTCGGCCAAGGGACCAAGGTGGAGATCAAAC	
mo:	e Kappa J region	
11100	41C	
JK 1		
JK2	TACACGTTCGGAGGGGGGCCACCAAGCTGGAAATAAAAC	
	TTCACATTCAGTGATGGGACCAGACTGGAAATAAAAC	
JK4	TICACHTICAGIGATOGGACCAGACTGGAAATAAAAC	
JK5		
Con:	ensus TTCGGTGGGGGACCAAGCTGGAAATAAAAC	
UIG	MJK] TGGTTCGACCTTTATTTTG,	
hum:	n Lambda pseudo J region	
	J1 C	
JPS	1 CACATGTTTGGCAGCAAGACCCAGCCCACTGTCTTAG	
mou:	e Lambda J region	
	JIC	
JL 1	TGGGTGTTCGGTGGAGGAACCAAACTGACTGTCCTAG	
JL2	TATGTTTTCGGCGGTGGAACCAAGGTCACTGTCCTAG	
JL3	TTTATTTTCGGCAGTGGAACCAAGGTCACTGTCCTAG	
	TECCCCCCCCCCA ACCA ACCA ACCACA	

FIG.2





A. SYNTHESIS OF HUMAN 1961 GENES a HUMAN 1961 HEAVY CHAIN STRUCTURE

V	DJ C	н1 н	CH2	снз	3'UT
	BstEII Apa I	Nar I	Loos-	-Smal -Hinf1	Smal
b. cDNA CLO	VES				100b
pGMH-3				Rsal Smal Hinfl	Rsal
pGMH-15	-BSIEII -Apo I	-Nar I -Bst E II - HinfI	-SacII	-Smo [-HinfI	-Smo!
pGMH-6	-BarE II -Apa J	-Nor I -BsrE II -Hinfl	Soci	-Smo I -Hinf!	-SmaI

B. A HUMAN IgG1 CONSTANT REGION CLONING VECTOR FOR V REGION MODULE INSERTION

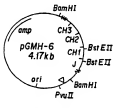
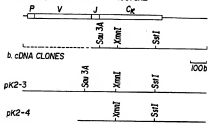
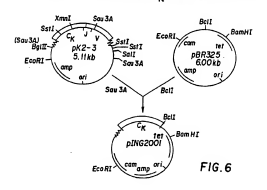


FIG.4

A. SYNTHESIS OF HUMAN IGK GENES a.HUMAN IGK LIGHT CHAIN STRUCTURE



B. CONSTRUCTION OF A HUMAN CK REGION CLONING VECTOR



7

TACTTTGACTACTGGGGCCAAGGCACCACTCTCACAGTCTCCTCAG GAGAGTGTCAGAGGGG GAGAGTGTCAGACGAGTCGGT

[MJH2]

ACCACACACACAGACGTCGGT TCCCTGAGACCAGAG ACCACAGAGAGAGAG

> [MJH3-8STEII] [MJH-BSTEII(13)]

CCTGGTTTGCTTACTGGGGCCAAGGGACTCTGGTCACTGTCTGCAG

Primers Designed for Ig V Region Synthesis

ton
Reg
٦ -
5
Cha
Heavy
19
÷

	Mismatches JH1 JH2 JH3 JH4	
Lon — CCATC	Hism JH1 J	0
H1 Reg	2	12
Human 1961 pGHI-6 GGTCACCGTCTCTCAG CCTCCACAAGGCCCATC BS/EII Basion	ס מעלדסונס מנוס גנדששנט	Jai tactgetacttcgatgetctggggcgcagggcccagggggggggg
Human 1961 pGMH-6		Gtactt
n IgGl		TACTG
Humar		3H1 T

F16.7A

7

JH4 TACTATGCTATGGACTACTGGGGTCAAGGAACCTCAGTCACCGTCTCCTCAG (HJH4)

JKS

17 0 21 3

~

JK2 TACAÇGITCGGAGGGGGCCAAGCTGGAATAAAAC [JK2BGLI]
CCCTGGTTCGAC<u>CTGATT</u>

Bg/II 0

Ξ

22

JK4 ATCACGTTCGGCTCGGGACAAGTTGGAAATAAAAC |SJK4| GCAAGCCGCTGT |JK48GLII| GCCCTGTTCAACTCTAAAAT JKS CTCACGTTCGGTGCTGGGACCAAGCTGGAGCTGAAAC [5JKS] GCAAGCCACGACCCTGG

Primers Designed for Ig V Region Synthesis

ion	CTGTCTTCATCTTCCC CTGTCTTCATCTTCCC		Hismatches JKl JK2 JK4	17 0 3 6
 H. Ig Kappa Chain J-C Region J Region Instant Region 	inman kappa pK1-3 CYGGAGATCAAAC GAACTGTGGCTGCATCTGGTTGATCTTCGC ping2001 $\frac{1GATGAAAC}{BGII}$	House kappa J Regions and Primers	2	JKI TGGACGTTCGGTGGAGGCACCAAGCTGGAATCAAAC (5JKI) GCAAGCCACCTCCGTGG

FIG.7A(cont.)

C. Mouse Variable Region Consensus Primers

AAGGTG GAGATGAAA ---- C - I------BstEII CCAGTGG CAGAG 0000 GGACCAC GGTCACC GTC GACC A'A GCTT GAG TTCGAA CT HINDIII TGTCAGAGGAGTCGGTCGTGTTT<u>CCCGGG</u>TA Apal 5° . U ,00000 700076533* ပ • 9 0 0 V 0000 4444 0000 9 9 9 0011000 944 72a J/C Junction Primer mouse heavy chain J segments mouse light chain J segments UIG-K consensus primer: VIG-H TACTGGTA TACTGGTA TACTATGC primer: JK1 JK2 JK4 JK5 consensus p D. House SCHL.60

54

HEAVY CHAIN V REGION MODULE GENE SYNTHESIS

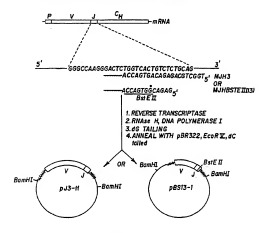
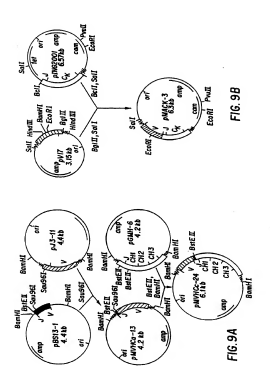
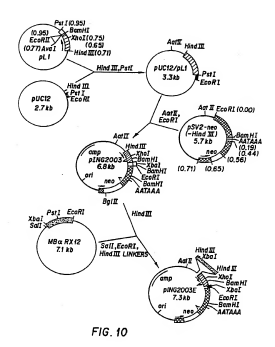


FIG. 8





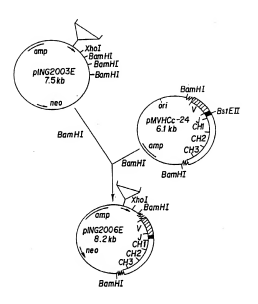


FIG. 11

Asp GAT 60	17 3er 7CA 120	37 Val GTC 180	Arg CCT 240	77 Ser AGC 300	Ala GCC 360	3er *	
Arg	Ala	177	GLY	Ser	107	Ser	
Ala	G1y GGG	CAC	Asa	Ser	TAC	Val S	
ACA TO	Pro	ATC	Ser	\$.5 \$.5	TAT	길입	
Ala	Lys	Tre	5 E	Asp C	Val 1 GTC 1	Val I	
Val GTA 45	Val 1 GTG 2	32 TYr 7 TAC 1	52 Asn E AAT (225	72 Val A GTA G	92 Ala V GCG G	## ## ## ## ## ## ## ## ## ## ## ## ##	
Ten	36 36	Ser 1	ATT A	Thr V	Ser A TCT G	Thr L	
Phe	C) n	ACC	CAG	Cle	Asp	613	
Te Cite	Ala	Phe	GCA GCA	P ACA	ggr Ggc	CAA	
Ile	G1y GGG	Acc	11e ATT	Ala	Ser	G1 y	
ATC 30	Pro CCT 90	Tyr Tyr TAC 150	47 Trp TCC 210	67 AAG 270	330 ACA	107 Tre 390	
Tyr	Gln	G13 GGC	Asp	Ser	Ge	TAC	
Ser							8
S S	Glu	Ser	35	Lys	Ser	Ala	2
17. 551	Leu 776	Ala	613	Phe	Ser	Phe TTT	FIG.12B
61 ,	Glu	Lys	CS.	Lys	ile.	17. 135	ш.
A TO A		22 Cys TGC 135	42 G1y GGA 195	62 Glu GAG 255	Gln CAA 315	102 Asp GAC 375	
ACC	CAG	Ser	Pro	Asn	APE	TAC	
rcc, ccc rct orc	Ser	Va 1	Arg	Tyr	TAC	Asp	
1, 1, 1, 1, 1, 1, 1, 1, 1, 1, 1, 1, 1, 1	His	Lys	GID	Asn	Ala GCC 1	Tyr 7	
GGA GGA		CTG	Lys AAG	ACT /	Thr A	Ser 1	
306E		. •				a F	

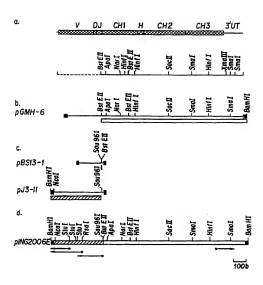


FIG.12A

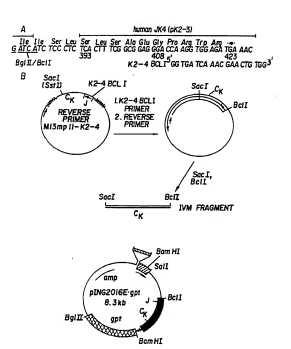


FIG. 13

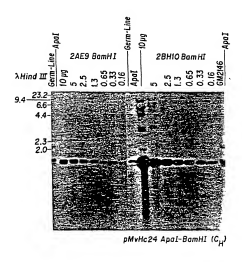
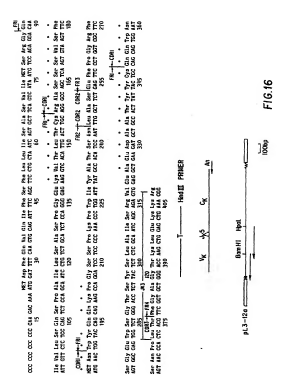


FIG. 14

358	75 CF 180	ATC 270	£ 5 8	GAC DA	£ 5 £		
AAC	F S	투 B	ξ£.	ظف	Asp GAT		
ξŞ	. ng 98	Lys AAG	A1a	122	66.9		
35	. 58	35	12.5	- 2 B	35		
ξā	. ភូម	657	F SC	15 F	Va 1	58	
Asp GAT 75	Lys 165	Lys Gly Leu L AAG GGT TTA A 255	Glu Thr Ser Ala T GAA ACC TCT GCC T 345	43 SE	525	Ser TCT 625	75
ATG ATG	CIN Ser Alba Can a The Cin Lea Val Cin Ser City Pro Ciu Lea Uye Lys Pro City City City Can Can Can Can Can City City City City City City City City	38	Ser Leu	ล้ย	A 20	Thr Trp Asn Ser ACC TGG AAC TCT 625	FIG. 15
ATC	35.	2 2	35	7 7 88	3 2	£g	Œ
GAC	G G G	A1a GCT	# 5 #	문학원	28	F SS	Пæл
Ĕ	· £5_	CAG CAG	Ala GCC	i ê E	7.1	32	[] SO]
56	• 982€	GLY Thr Phe Thr Jan Tyr GLY BET AsnTrp Val Lys Gln Ala Pro GGG TAT ACC TTC ACA ARC TAT GGA ATG AAC TGG GGG AGG CGG GCT CCA . 210 22 240	Asp Phe Lys Gly Arg Phe Ala Phe GAC TTC And GOA CGG TTT GCC TTC 315	2558	STC STC	Thr Leu Thr T ACC TTG ACC T 600	4
CAA	. P. T.	75 4a1	ya Gly Arg P	48	75	35	
8	55	7 Tro	90,5	45	£ 5	2.5	
72	14 SP	NS A	85 E	£Ε	4 S	98	
CAC	38 3	DA MG	ěΕ	žΣ	ĘŽ	£ 5	
CAC 45	955	38.5°	Asp GAC 315	F & 50	₹ 5 £	55 T 28	25
Ş	Te TE	\$\$	GAT	45	Lys AA	ŽĮ.	тншов.
Ē	188	A Sa	S P	ATG	4 S	86	(
55	A La	1 2	Gly Gln Pro Thr Tyr Ala Asp GGA CAG CCA ACA TAT GCT GAT 300	GAC	r Val Ser Ser Al	The Leu Cly Cys Leu Val Lys Cly Tyr Phe Pro Clu Pro Act cta GGA TGC CTG GTC AAG GGT TAT TTC CCT GAG CCA 1 570 570	F = 100 H
E	ga a	£ £	έŞ	e e e	7 8 5	Va 1 CTC	≥ II108-14
30 06	APR 60 120	4 50 5 50 50	5 2 8	AAT 390	12 E 8	356	IHmo8
ວິ	385	žž	CAG CAG	Lys	ĘŞ	35	.6a Fi
8	5 5	9,99	58	35	15 15	28	3-6
9	Ala	à t	Tyr Thr TAC ACT	ASD	وَعَ	35	_
8	Ala Ala Ala G GCA GCT GCC C 105	Cys Lys Ala Ser TCC AAG GCT TCT 195	7.5	AAC	ĘŞ	ĘŞ	
25 25	A SCA	AAG 195	14 S	11e ATC 375	£88.8	Ser Val 1 TCG GTG 1 555	
22	ATC A	\$ 55 20 20 20 20 20 20 20 20 20 20 20 20 20	AAC	CAG CAG	CAA CAA	\$ 55	
60 APC CCC CCC CCC CCC CCC CAC TTT GTC TTA ACC CAC CAC TCA CCC CAA CTC TTA GLC ATC ATG GAT TGG CTC 15	CTA TTC CTG /	AAG ATC TCC	GG TG ATA AAC AC	Als by Leu Gin lie fan kan Leu Lys kan Giu kap Net Als The Try Pae Cys als Alfelfne Ser by. Giy kan Ser her lyp Ser GCC Tht THO GIG ATE MA ALS GEG ALA ALS GOOD ALS AND THE TOT GCA AIGHT THO CON THE TOT GCT TO THO THE TOT TOT GCT AIGHT AND ALS AND	2014—150 day the The Lea The Val Ser Ser all 159 The The Ala Poo Ser Val Pre Poo Lea Ala Poo Val Ogs Cly App The Tho Coc Cas Ogs Loc Act one Lea The Test Good Ala Cas And Good Cas The Cas Off The Cas one Good Coc One The Cas Act The Tho Coc Cas Ogs Loc Act one Lea The Valo	ACT GGC TCC 1	
ATC.	4.5	- E	TE 5	Ç.	I E B	G13	
8	35	2 8 8 B	458	A) a	중주동	μŞ	

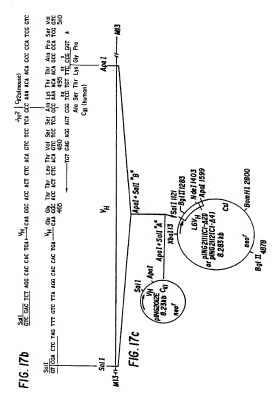


 $\frac{BanH}{4}$ we have the case case case case case that the case case case and $\frac{BanH}{32}$ whereas $\frac{BanH}{32}$ SOLT GIC GAC TET AGG CAC CAC TGA GCC CAA GTC TTA GAC ATG GAT TGG CTG TGG AAC TTG

SALI GT CGA CTC TAG TIT GTC TTA AGG CAC CAC TGA GCC CAA GTC TTA GAC ATG GAT TGG CTG TGG AAC TTG

F16. 17a

CI-421 CI-44



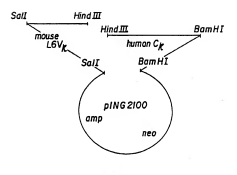
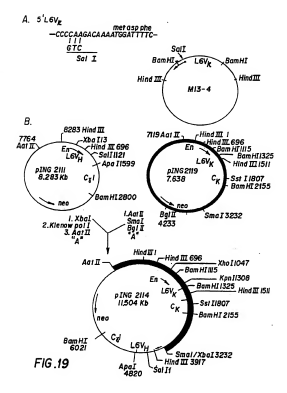


FIG.18



œ	
6	
ũ	
•	:
Ω	

V_H pH3-64 (J_H2) oligo (dT) clone, BAL-31 delections 5'. CXI APA mutagenesis—pING 2111 neo

PING2112

pinG2III

GTCGACTCTAGTTTGTCTTAAGGCACCACTGAGCCCAAG TCTTAGACAT CATGGAT

ACC ACT CTC ACA GTC TCC TCA GCC AGC ACA AAG GGC molhu Joint

 V_{K} p13–12A (J $_{K}$ 5) oligo (dT) clone. J $_{K}$ Hind III mulagenesis, 5 $^{\circ}$ SAL mulagenesis — pING 2120 got p1NG 2120 got

Soll met 5' GTC GAC AAA ATG GAT

K joint accade ct(T)gag(A)TG aaa|cga act

FIG. 20

2HT heavy chain variable sequence met gly phe ser arg lie phe control of the con

FIG. 21

2H7 light chain variable sequence

met aap phe gin val gin iie phe ser phe leu leu

C23CCCAAAATCAAAGCAAAATG GAT TTT CAA CTG CAG ATT TTC AGC TTC CTG CTA

Soil pumer Fil

ile ser ala ser val iie ile ala arg gly gin ile val leu ser gin ser
ATC AGT GCT TCA GTC ATA ATT GCC AGA GAG CAA ATT GTT CCT CTC CAG TCT

pro ala ile leu ser ala ser pro gly glu lys val thr met thr cys arg cCC AGC ATC CTG TCT GCC AGT TCC AGG GCG AGA GAG CAC ATT GTT CCA GG

CCR!

ala ser ser ser val ser tyr Cmg! FR2

ala ser ser ser val ser tyr Cmg! FR2

cCC AGC TCA AGT TTA AGT TAC ATT GCA CTG TAC CAG CAG AGA GCC AGG TCC

FR2

Ser pro lys pro trp ile tyr lala pro ser asi leu ala ser gily val pro
TCC CCC AAA CCC TGG ATT TAT GCC CCA CAC CAC CAG CAG CAG CCC

ala arg phe ser gly ser gly ser gly thr ser tyr en leu thr ile ser
CCT CGC TTC AGT GGG AGT GGG TCT GGG ACC TCT TAC TCT CAC ATC AGC

arg val glu ala glu asp ala ala thr tyr tyr cys gin gin trp ser phe
AGG GTG GAG GGT GAG GAT GCG CCC ACC TAT TAT TAC TCC CCC

. CDR3, FF4

asn pro pro thr phe gly ala gly thr lys leu glu leu Tys
AAC CCA CCC ACG TCT GCT GCG ACC AGC AGA GCC AGG TTT

ACC CAG TCC GCT GCT GCT GCG ACC AGG CTG AAA

CCC CCC CCC AGT TC GCT GCT GCG ACC AGG CTG AAA

CCC CCC CCC AGT TC GCT GCT GCG ACC AGC AGC CCA CAG

ACC CCAC CCC CCT GCT GCT GCG ACC AGC AGC CTG AAA

CCC CCC CCC AGT TC GCT GCT GCG ACC AGC AGC CTG AAA

CCC CCC CCC AGC TC GCT GCT GCG ACC AGC AGC CTG AAA

CCC CCC CCC AGC TC GCT GCT GCG ACC AGC AGC CTG AAA

CCC CCC CCC AGC TC CGT GCT GCG ACC AGC AGC CTG AAA

CCC CCC CCC AGC TC CGT GCT GCG ACC AGC AGC CTG AAA

CCC CCC CCC CCC AGC TC CGT GCT GCG ACC AGC AGC CTG AAA

CCC CCC CCC AGC TC CGT GCT GCG ACC AGC ACC AGC CTG AAA

CCC CCC CCC CCC ACC TCC TCC CCC ACC AGC CCC ACC

TCC GCC AAAC CCC CCC ACC TCC ACC

TCC GCC AACC CCC ACC TCC ACC

TCC GCC ACC TCC ACC

TCC ACC ACC ACC

TCC ACC ACC ACC

TCC ACC

TCC ACC

TCC ACC

TCC ACC

TCC A

JKHIND III primer

FIG. 22

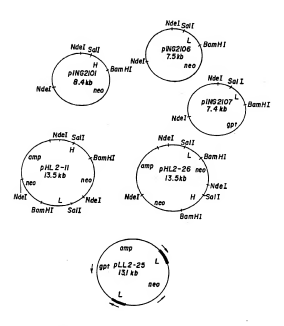


FIG. 23

2H7 Chimerae

V_H pH2-7 (J_H1) J_HBSIEII clone, NGOI cut 5'ATG —pING2101 neo

Sall met TCGACATGGGA

5' **6**TC 6A<u>CATG 66A</u> mo_l hu

mo, hu Cvi

p. $2-i2(i_{\rm H}5)$ oligold 7 clone, $J_{\rm K}$ Hind $I\!\!II$ mutagenesis, 5 'SAL mutagenesis—piNG2105 gpr pR-2207 gpr

Sals met † GTC GAC AAA ATG GAT neg hu acc ade ct De a @ De aaa ce

Joint

F16. 24

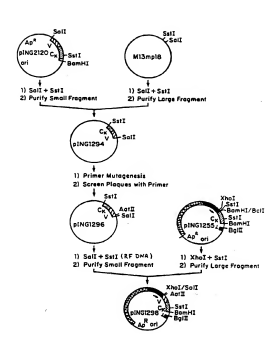


FIG.25A

5' ATA ATG TCC AGA CGT CAA ATT GTT 3'

Aota

Signal Sequence Processing Site

AIG GAT TIT CAA GTG CAG ATT TTC AGC TTC CTG CTA ATC ACT TCA GTC ATA ATG TCC AGA GGA CAA ATT GTT CTC TCC CAG TCT CCA GCA MET hap the Gin Val Gin lie the Ser the Leu Leu lie Ser Ala Ser Val lie MET Ser Arg Gly Gin lie Val Leu Ser Gin Ser Pro Ala

FIG. 25B

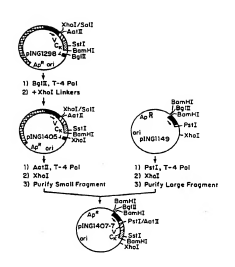


FIG.25C

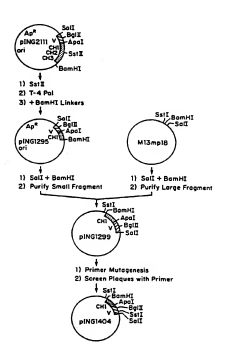


FIG.26A

5' AM AGT GCC CGA GCT CAG ATC CAG TTG GT 3'

Signal Sequence Processing Site MET ASP Trp Leu Trp Asn Leu Leu Phe Leu MET Ala Ala Ala Gin Ser Ala Gin Ala Gin Ile Gin Leu Val Gin Ser Gly Pro Glu ATG GAT TGG CTG TGG AAC TTG CTA TTC CTG ATG GCA GCT GCC CAA AGT GCC CAA GCA CAG ATC CAG TTG GTG CAG TCT GGA CCT GAG

FIG.26B

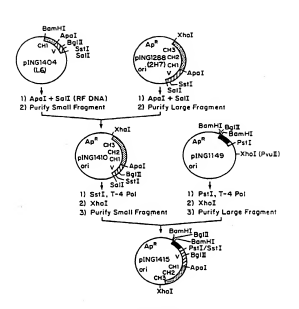
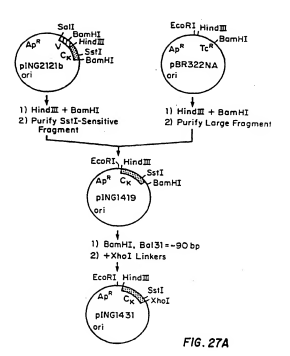
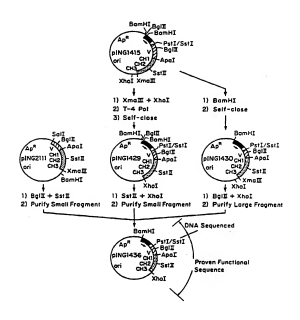
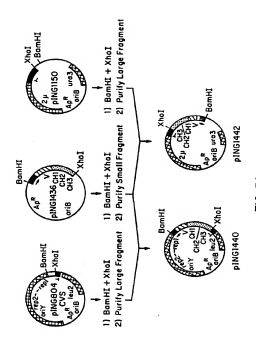


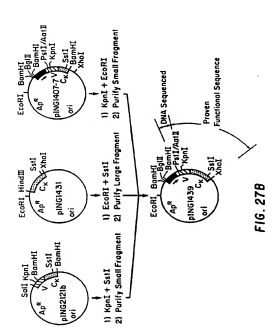
FIG.26C

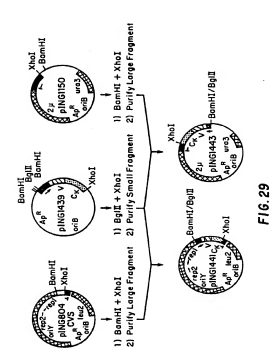




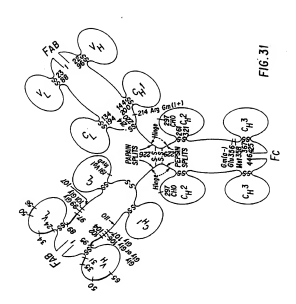
F1G.28

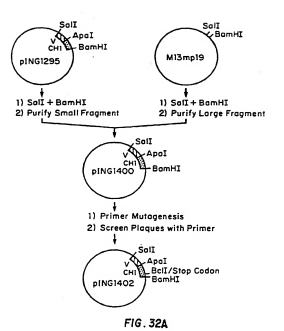




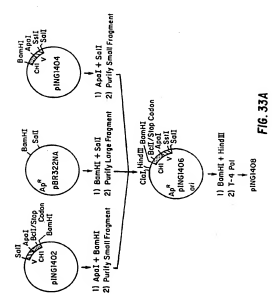


84





86



Stop Codon



FIG. 32B

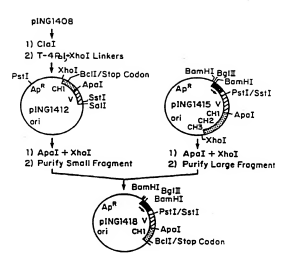


FIG. 33B

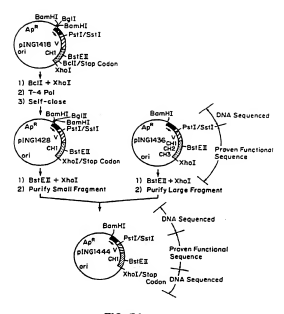


FIG. 34

Dral

TIT AAA AGG AAA TTT TIT CTT ATA AAA

CCC AAA TTA TCC AAT CAT CAG TAT TAC AAA ATG TTT CAA CCG TAA TAC ATT TAA CAT TTC

ACC CTT GAA CTG ATC TTA TTT TTT GAC CAC ACT CCC CTT GGT TTT TCA CCA AAN CTG AGT

Nde I

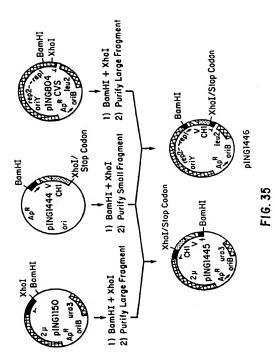
TTC ATT TTT GTT GAA AAA TTT GTA CCT GCG ACA TCG GGC ATA TGG AAC GAT AAA TGC CCA

1 MET Lys Tyr Leu Leu Pro Thr Ala Ala Ala TGA AMA TTC TAT TTC AMG GAG ACA GTC ATA ATG AMA TAC CTA TTG CCT ACG GCA GCC GCT

Hge III

Gly Leu Leu Leu Ala Ala Gln Pro Ala MET Ala Ala Asn Thr Gly Gly Tyr Ala Thr GGA TTG TTA TTA CTC GCT GCC CAA CCA GCG ATG GCC GCA AAT ACG GGT GGC TAT GCC ACC

FIG. 36 A



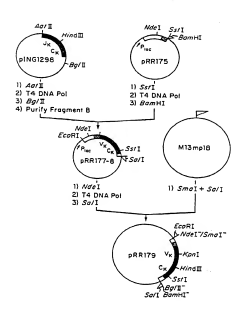


FIG. 37A

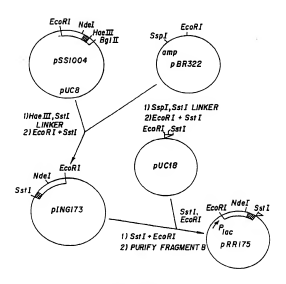


FIG. 36B

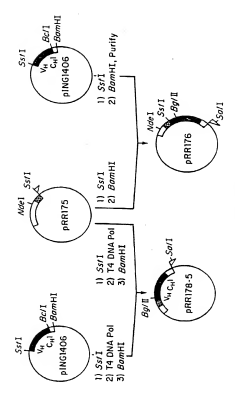
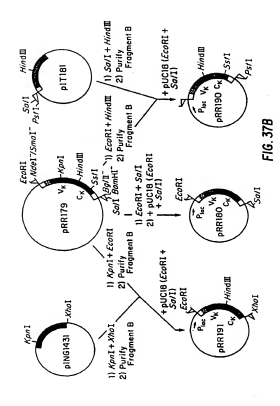


FIG.38A



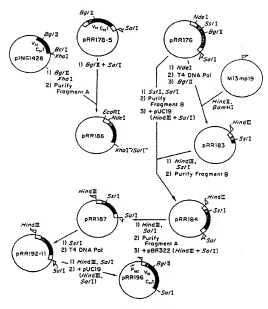


FIG. 38B

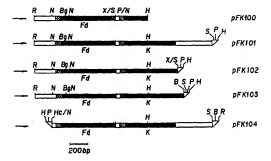
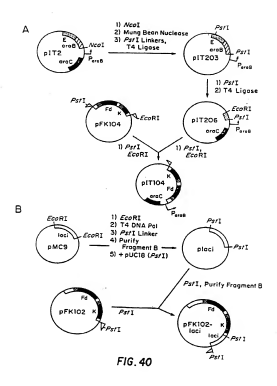


FIG. 39



This Page is Inserted by IFW Indexing and Scanning Operations and is not part of the Official Record

BEST AVAILABLE IMAGES

Defective images within this document are accurate representations of the original documents submitted by the applicant.

Defects in the images include but are not limited to the items checked:

☐ BLACK BORDERS	
\square image cut off at top, bottom or sides	
☐ FADED TEXT OR DRAWING	
☐ BLURRED OR ILLEGIBLE TEXT OR DRAWING	
☐ SKEWED/SLANTED IMAGES	
\square color or black and white photographs	
☐ GRAY SCALE DOCUMENTS	
☐ LINES OR MARKS ON ORIGINAL DOCUMENT	
☐ REFERENCE(S) OR EXHIBIT(S) SUBMITTED ARE POOR QUALITY	

IMAGES ARE BEST AVAILABLE COPY.

As rescanning these documents will not correct the image problems checked, please do not report these problems to the IFW Image Problem Mailbox.